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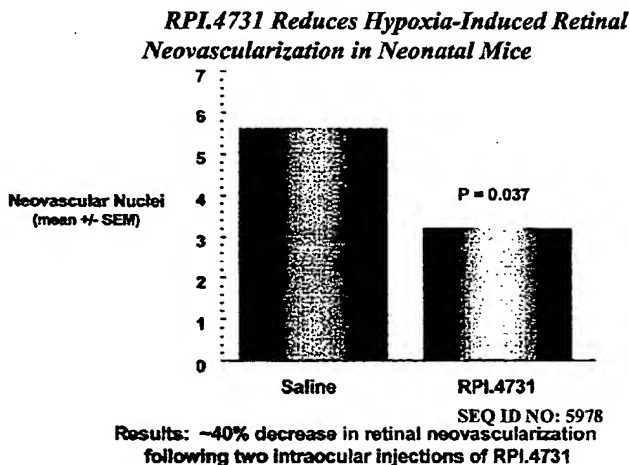
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- (57) Abstract: The present invention relates to nucleic acid molecules, including dsRNA, siRNA, antisense, 2,5-A chimeras, aptamers, and enzymatic nucleic acid molecules, such as hammerhead ribozymes, DNAzymes, and allozymes, which modulate the expression of vascular endothelial growth factor receptor (VEGF) and/or vascular endothelial growth factor receptor (VEGFr) genes for the treatment and/or diagnosis of diseases and conditions associated with angiogenesis, such as cancer, tumor angiogenesis, or ocular indications such as diabetic retinopathy, or age related macular degeneration, proliferative diabetic retinopathy, hypoxia-induced angiogenesis, rheumatoid arthritis, psoriasis, wound healing, and female reproductive disorders and conditions, including but not limited to endometriosis, endometrial carcinoma, gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), and menopausal dysfunction.



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NUCLEIC ACID BASED MODULATION OF FEMALE REPRODUCTIVE DISEASES  
AND CONDITIONS

This patent application claims priority from Sandberg *et al.*, USSN 60/334,461, filed November 30, 2001, entitled "Method and Reagent for the Modulation of Female Reproductive Diseases and Conditions" and Pavco *et al.*, USSN 10/138,674, filed May 3, 2002, which is a continuation in part of Pavco *et al.*, USSN 09/870,161, which is a continuation-in-part of Pavco *et al.*, USSN 09/708,690, filed November 7, 2000, which is a continuation-in-part of Pavco *et al.*, USSN 09/371,722, filed August 10, 1999, which is a continuation-in-part of Pavco *et al.*, USSN 08/584,040, filed January 11, 1996, which claims the benefit of Pavco *et al.*, USSN 60/005,974, filed on October 26, 1995; these earlier applications are entitled "Method and Reagent for Treatment of Diseases or Conditions Related to Levels of Vascular Endothelial Growth Factor Receptor". Each of these applications is hereby incorporated by reference herein in its entirety including the drawings and tables.

Technical Field Of The Invention

This invention relates to methods and reagents for the treatment of diseases or conditions relating to the levels of expression of vascular endothelial growth factor (VEGF) and vascular endothelial growth factor receptor(s). Specifically, the instant invention features nucleic-acid based molecules and methods that modulate the expression of vascular endothelial growth factor and/or vascular endothelial growth factor receptors, such as VEGFR1 and/or VEGFR2, that are useful in preventing, treating, controlling and/or diagnosing disorders and conditions related to angiogenesis, including but not limited to cancer, tumor angiogenesis, or ocular indications such as diabetic retinopathy, or age related macular degeneration, proliferative diabetic retinopathy, hypoxia-induced angiogenesis, rheumatoid arthritis, psoriasis, wound healing, endometriosis, endometrial carcinoma, gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), and menopausal dysfunction.

Background Of The Invention

The following is a discussion of relevant art, none of which is admitted to be prior art to the present invention.

VEGF, also referred to as vascular permeability factor (VPF) and vasculotropin, is a potent and highly specific mitogen of vascular endothelial cells (for a review see Ferrara, 1993 *Trends Cardiovas. Med.* 3, 244; Neufeld *et al.*, 1994, *Prog. Growth Factor Res.* 5, 89). VEGF-induced neovascularization is implicated in various pathological conditions such as  
5 tumor angiogenesis, or ocular indications such as diabetic retinopathy, or age related macular degeneration, proliferative diabetic retinopathy, hypoxia-induced angiogenesis, rheumatoid arthritis, psoriasis, wound healing and others.

VEGF, an endothelial cell-specific mitogen, is a 34-45 kDa glycoprotein with a wide range of activities that include promotion of angiogenesis, enhancement of vascular-  
10 permeability and others. VEGF belongs to the platelet-derived growth factor (PDGF) family of growth factors with approximately 18% homology with the A and B chain of PDGF at the amino acid level. Additionally, VEGF contains the eight conserved cysteine residues common to all growth factors belonging to the PDGF family (Neufeld *et al.*, *supra*). VEGF protein is believed to exist predominantly as disulfide-linked homodimers; monomers of  
15 VEGF have been shown to be inactive (Plouet *et al.*, 1989 *EMBO J.* 8, 3801).

VEGF exerts its influence on vascular endothelial cells by binding to specific high-affinity cell surface receptors. Covalent cross-linking experiments with  $^{125}$ I-labeled VEGF protein have led to the identification of three high molecular weight complexes of 225, 195 and 175 kDa presumed to be VEGF and VEGF receptor complexes (Vaisman *et al.*, 1990 *J. Biol. Chem.* 265, 19461). Based on these studies VEGF-specific receptors of 180, 150 and 130 kDa molecular mass were predicted. In endothelial cells, receptors of 150 and 130 kDa have been identified. The VEGF receptors belong to the superfamily of receptor tyrosine kinases (RTKs) characterized by a conserved cytoplasmic catalytic kinase domain and a hydrophilic kinase sequence. The extracellular domains of the VEGF receptors consist of  
20 seven immunoglobulin-like domains that are thought to be involved in VEGF binding functions.  
25

The two most abundant and high-affinity receptors of VEGF are flt-1 (VEGFR1) (*fms*-like tyrosine kinase) cloned by Shibuya *et al.*, 1990 *Oncogene* 5, 519 and KDR (VEGFR2) (kinase-insert-domain-containing receptor) cloned by Terman *et al.*, 1991 *Oncogene* 6, 1677.  
30 The murine homolog of KDR, cloned by Mathews *et al.*, 1991, *Proc. Natl. Acad. Sci., USA*, 88, 9026, shares 85% amino acid homology with KDR and is termed as flk-1 (fetal liver kinase-1). The high-affinity binding of VEGF to its receptors is modulated by cell surface-associated heparin and heparin-like molecules (Gitay-Goren *et al.*, 1992 *J. Biol. Chem.* 267, 6093).

VEGF expression has been associated with several pathological states such as tumor angiogenesis, several forms of blindness, rheumatoid arthritis, psoriasis and others. In addition, a number of studies have demonstrated that VEGF is both necessary and sufficient for neovascularization. Takashita *et al.*, 1995 *J. Clin. Invest.* 93, 662, demonstrated that a 5 single injection of VEGF augmented collateral vessel development in a rabbit model of ischemia. VEGF also can induce neovascularization when injected into the cornea. Expression of the VEGF gene in CHO cells is sufficient to confer tumorigenic potential to the cells. Kim *et al.*, *supra* and Millauer *et al.*, *supra* used monoclonal antibodies against VEGF or a dominant negative form of VEGFR2 receptor to inhibit tumor-induced 10 neovascularization.

During development, VEGF and its receptors are associated with regions of new vascular growth (Millauer *et al.*, 1993 *Cell* 72, 835; Shalaby *et al.*, 1993 *J. Clin. Invest.* 91, 2235). Furthermore, transgenic mice lacking either of the VEGF receptors are defective in blood vessel formation and these mice do not survive; VEGFR2 appears to be required for 15 differentiation of endothelial cells, while VEGFR1 appears to be required at later stages of vessel formation (Shalaby *et al.*, 1995 *Nature* 376, 62; Fung *et al.*, 1995 *Nature* 376, 66). Thus, these receptors apparently need to be present to properly signal endothelial cells or their precursors to respond to vascularization-promoting stimuli.

Increasing evidence suggests that the VEGF family may also be involved with both the 20 etiology and maintenance of peritoneal endometriosis. Peritoneal endometriosis is a significant debilitating gynecological problem of widespread prevalence. It is now generally accepted that the pathogenesis of peritoneal endometriosis involves the implantation of exfoliated endometrium. Maintenance of exfoliated endometrial tissue is dependent upon the generation and maintenance of an extensive blood supply both within and surrounding the 25 ectopic tissue.

Endometriosis is a disease affecting an estimated 77 million women and teenagers worldwide. Endometriosis is a leading cause of infertility, chronic pelvic pain and hysterectomy. Endometriosis can be characterized when endometrial tissue (the tissue inside the uterus which builds up and is shed each month during menses) is found outside the uterus, 30 in other areas of the body. The endometrial tissue can respond to hormonal commands each month and break down and bleed. However, unlike the endometrium, these tissue deposits have no way of leaving the body. The result is internal bleeding, degeneration of blood and tissue shed from the growths, inflammation of the surrounding areas, expression of irritating enzymes and formation of scar tissue. In addition, depending on the location of the growths,

interference with the bowel, bladder, intestines and other areas of the pelvic cavity can occur. Endometrial tissue has even been found lodged in the skin and at other extrapelvic locations like the arm, leg and even brain.

Currently, the presence of Endometriosis can only be confirmed through surgery such  
5 as laparoscopy, but can be suspected based on symptoms, physical findings and diagnostic tests. Endometriosis can be treated in many different ways, both surgically and medically. Most commonly, surgery will be performed during which the disease will be excised, ablated, fulgurated, cauterized or otherwise removed, and adhesions will also be freed. Surgeries include but are not limited to laparoscopy; laparotomy; presacral and uterosacral and various  
10 levels of hysterectomies, where some or all of the reproductive organs are removed. Often, this method will only relieve the symptoms associated with growths on the reproductive organs, not the bowels or kidneys and related areas where Endometriosis can be present.

There are several drugs used to treat Endometriosis that are utilized either alone or in combination with surgery. These include contraceptives, GnRH agonists, and/or synthetic  
15 hormones. GnRH agonists are commonly used on women in all stages of the disease and may sometimes have serious side affects. GnRH (gonadotropin releasing hormone) analogues are classified into 2 groups: agonists and antagonists. Agonists are commonly used in the treatment of Endometriosis by suppressing the manufacture of follicle stimulating hormone (FSH) and luteinizing hormone (LH), common hormones required in ovulation. When they  
20 are not secreted, the body will go into "pseudo-menopause," stalling the growth of more implants. However, these are again only stop-gap measures that can be utilized only for short term intervals. Once the body returns to it's normal state, the Endometriosis will again begin to implant itself.

Angiogenesis is likely to be involved in the pathogenesis of endometriosis. According  
25 to the transplantation theory, when the exfoliated endometrium is attached to the peritoneal layer, the establishment of a new blood supply is essential for the survival of the endometrial implant and development of endometriosis (Donnez *et al.*, 1998, *Hum. Reprod.*, 13, 1686-1690). Endometrial growth and repair after menstruation are associated with profound angiogenesis. Abnormalities in these processes result in excessive or unpredictable bleeding  
30 patterns and are common in many women. It is therefore important to understand which factors regulate normal endometrial angiogenesis. Vascular endothelial growth factor (VEGF) is an endothelial cell-specific mitogen that plays an important role in normal and pathological angiogenesis (Fasciani *et al.*, 2000, *Mol. Hum. Reprod.*, 6, 50-54; Sharkey *et al.*, 2000, *J. Clin. Endocrinol. Metab.*, 85, 402-409). Sources of this factor include the eutopic

endometrium, ectopic endometriotic tissue and peritoneal fluid macrophages. Important to its etiology is the correct peritoneal environment in which the exfoliated endometrium is seeded and implants. Established ectopic tissue is then dependent on the peritoneal environment for its survival, an environment that supports angiogenesis. The increasing knowledge of the 5 involvement of the VEGF family in endometriotic angiogenesis raises the possibility of novel approaches to its medical management, with particular focus on the anti-angiogenic control of the action of VEGF (McLaren, 2001, *Hum. Reprod. Update*, 6, 45-55).

10 Pavco *et al.*, International PCT Publication No. WO 97/15662, describes methods and reagents for treating diseases or conditions related to levels of vascular endothelial growth factor receptor.

Robinson, International PCT Publication No. WO 95/04142, describes the use of certain antisense oligonucleotides targeted against VEGF RNA to inhibit VEGF expression.

Jelinek *et al.*, 1994 *Biochemistry* 33, 10450 describe the use of specific VEGF-specific high-affinity RNA aptamers to inhibit the binding of VEGF to its receptors.

15 Rockwell and Goldstein, International PCT Publication No. WO 95/21868, describe the use of certain anti-VEGF receptor monoclonal antibodies to neutralize the effect of VEGF on endothelial cells.

20 Pappa, International PCT Publication No. WO 01/32920, describes inhibitors, including certain ribozyme and antisense nucleic acid molecules, of specific genes, including cathepsin D, AEBP-1, stromelysin-3, cystatin B, protease inhibitor 1, sFRP4, gelsolin, IGFBP-3, dual specificity phosphatase 1, PAEP, Ig gamma chain, ferritin, complement component 3, pro-alpha-1 type III collagen, proline 4-hydroxylase, alpha-2 type I collagen, claudin-4, melanoma adhesion protein, procollagen C-endopeptidase enhancer, nascent-polypeptide-associated complex alpha polypeptide, elongation factor 1 alpha (EF-1-alpha), vitamin D3 25 hydroxylase, CSRP-1, steroidogenic acute regulatory protein, apolipoprotein E, transcobalamin II, prosaposin, early growth response 1 (EGR1), ribosomal protein S6, adenosine deaminase RNA-specific protein, RAD21, guanine nucleotide binding protein beta polypeptide 2-like 1 (RACK1) and podocalyxin genes which are all differentially expressed in tissues within individual patients with endometriosis.

30 Labarbera *et al.*, International PCT Publication No. WO 00/73416, describes specific antisense nucleic acid molecules targeting follicle-stimulating hormone receptor.

Storella *et al.*, International PCT Publication No. WO 99/63116, describes modulators of Prothymosin gene products for treating endometriosis, including certain ribozymes and antisense nucleic acid molecules.

#### Summary Of The Invention

5

This invention features nucleic acid-based molecules, for example, enzymatic nucleic acid molecules, allozymes, antisense nucleic acids, 2-5A antisense chimeras, triplex forming oligonucleotides, decoy RNA, dsRNA, siRNA, aptamers, and antisense nucleic acids containing nucleic acid cleaving chemical groups, and methods to modulate vascular endothelial growth factor (VEGF) and/or vascular endothelial growth factor receptor (VEGFr) gene expression. Non-limiting examples of genes that encode vascular endothelial growth factor receptors of the invention include VEGFR1, VEGFR2 or combinations thereof. In particular, the instant invention features nucleic acid-based molecules and methods that modulate the expression of vascular endothelial growth factor and/or vascular endothelial growth factor receptors, such as VEGFR1 and/or VEGFR2, that are useful in preventing, treating, controlling, and/or diagnosing angiogenesis related diseases and conditions, including but not limited to tumor angiogenesis, cancers such as breast cancer, lung cancer, colorectal cancer, renal cancer, pancreatic cancer, or melanoma, or ocular indications such as diabetic retinopathy, or age related macular degeneration, and female reproductive disorders and conditions, including but not limited to endometriosis, endometrial carcinoma, gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), and menopausal dysfunction.

In one embodiment, the invention features one or more nucleic acid-based molecules and methods that independently or in combination modulate the expression of gene(s) encoding vascular endothelial growth factor receptors. Specifically, the present invention features nucleic acid molecules that modulate the expression of VEGF (for example Genbank Accession No. NM\_003376), VEGFR1 receptor (for example Genbank Accession No. NM\_002019), and VEGFR2 receptor (for example Genbank Accession No. NM\_002253) that are useful in preventing, treating, controlling, and/or diagnosing tumor angiogenesis, cancers such as breast cancer, lung cancer, colorectal cancer, renal cancer, pancreatic cancer, or melanoma, or ocular indications such as diabetic retinopathy, or age related macular degeneration, and female reproductive disorders and conditions, including but not limited to

endometriosis, endometrial carcinoma, gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), and menopausal dysfunction.

In one embodiment, the present invention features a compound having Formula I: (SEQ ID NO: 5977)



wherein each a is 2'-O-methyl adenosine nucleotide, each g is a 2'-O-methyl guanosine nucleotide, each c is a 2'-O-methyl cytidine nucleotide, each u is a 2'-O-methyl uridine nucleotide, each A is adenosine, each G is guanosine, each s individually represents a phosphorothioate internucleotide linkage, U is 2'-deoxy-2'-C-allyl uridine, and B is an inverted deoxyabasic moiety. This compound is also referred to as ANGIOZYME™ ribozyme.

In another embodiment, the present invention features a compound having Formula II: (SEQ ID NO: 5978).



15

wherein each a is 2'-O-methyl adenosine nucleotide, each g is a 2'-O-methyl guanosine nucleotide, each c is a 2'-O-methyl cytidine nucleotide, each u is a 2'-O-methyl uridine nucleotide, each A is adenosine, each G is guanosine, each s individually represents a phosphorothioate internucleotide linkage, U is 2'-deoxy-2'-C-allyl uridine, and B is an inverted deoxyabasic moiety.

In one embodiment, the invention features a composition comprising a nucleic acid molecule of the invention in a pharmaceutically acceptable carrier. In another embodiment, the invention features a composition comprising a compound of Formula I and/or Formula II in a pharmaceutically acceptable carrier or diluent.

25

In one embodiment, the invention features a method of administering to a cell, for example a mammalian cell, including a human cell, a nucleic acid molecule of the invention comprising contacting the cell with the nucleic acid molecule under conditions suitable for administration, for example in the presence of a delivery reagent such as a lipid, cationic lipid, phospholipid, or liposome. In another embodiment, the invention features a method of 30 administering to a cell, for example a mammalian cell, including a human cell, a compound of Formula I and/or Formula II comprising contacting the cell with the compound under

conditions suitable for administration, for example in the presence of a delivery reagent such as a lipid, cationic lipid, phospholipid, or liposome.

In one embodiment, the present invention features a mammalian cell comprising a nucleic acid molecule of the invention, wherein the mammalian cell is, for example, a human cell. In another embodiment, the present invention also features a mammalian cell comprising the compound of Formula I and/or Formula II, wherein the mammalian cell is, for example, a human cell.

In one embodiment, the invention features a method of inhibiting angiogenesis, for example tumor angiogenesis, or ocular indications such as diabetic retinopathy, or age related macular degeneration, or endometrial neovascularization, in a subject comprising contacting the subject with a nucleic acid molecule of the invention, under conditions suitable for the inhibition. In another embodiment, the invention features a method of inhibiting angiogenesis, for example tumor angiogenesis, or ocular indications such as diabetic retinopathy, or age related macular degeneration, or endometrial neovascularization, in a subject, comprising contacting the subject with a compound of Formula I and/or Formula II, under conditions suitable for the inhibition.

In another embodiment, the invention features a method of treatment of a subject having an ocular condition associated with the increased level of a VEGF receptor, for example diabetic retinopathy, or age related macular degeneration, comprising contacting cells of the subject with a nucleic acid molecule, such as an enzymatic nucleic acid molecule targeted against a VEGF receptor RNA, e.g., molecule according to Formula I and/or II, under conditions suitable for the treatment.

In another embodiment, the invention features a method of treatment of a subject having a condition associated with an increased level of VEGFR and/or a VEGF receptor, for example tumor angiogenesis, cancers such as breast cancer, lung cancer, colorectal cancer, renal cancer, pancreatic cancer, or melanoma, ocular diseases or ocular indications such as diabetic retinopathy, or age related macular degeneration, rheumatoid arthritis, psoriasis, endometriosis, endometrial carcinoma, gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), or menopausal dysfunction, comprising contacting cells of the subject with a nucleic acid molecule of the invention, such as a compound of Formula I and/or Formula II, under conditions suitable for the treatment.

In yet another embodiment, the inventive method of treatment further comprises the use of one or more drug therapies under conditions suitable for the treatment. Non-limiting

- examples of other drug therapies that can be used in combination with nucleic acid molecules of the invention include to 5-fluoro uridine, Leucovorin, Irinotecan (CAMPTOSAR® or CPT-11 or Camptothecin-11 or Campto), Paclitaxel, or Carboplatin, GnRH (gonadotropin releasing hormone) agonists, Lupron Depot (Leuprolide Acetate), Synarel (naferalin acetate),  
5 Zolodex (goserelin acetate), Suprefact (buserelin acetate), Danazol, or oral contraceptives including but not limited to Depo-Provera or Provera (medroxyprogesterone acetate), or any other estrogen/progesterone contraceptive.

In one embodiment, the invention features a method of administering to a mammal, for example a human, a nucleic acid molecule of the invention comprising contacting the mammal with the nucleic acid molecule under conditions suitable for the administration, for example, in the presence of a delivery reagent such as a lipid, cationic lipid, phospholipid, or liposome. In another embodiment, the invention features a method of administering to a mammal, for example a human, a compound of Formula I and/or Formula II comprising contacting the mammal with the compound under conditions suitable for the administration,  
10 for example, in the presence of a delivery reagent such as a lipid, cationic lipid, phospholipid, or liposome.  
15

In one embodiment, the invention features a nucleic acid molecule which down regulates expression of a vascular endothelial growth factor (VEGF) and/or vascular endothelial growth factor receptor (VEGFr) gene, for example, wherein the VEGFr gene  
20 comprises VEGFR1 or VEGFR2 and any combination thereof.

In one embodiment, a nucleic acid molecule of the invention, such as an enzymatic nucleic acid molecule, antisense nucleic acid molecule, 2-5A antisense chimera, triplex forming oligonucleotide, decoy RNA, dsRNA, siRNA, aptamer, or antisense nucleic acid containing nucleic acid cleaving chemical groups, is adapted to treat, control and/or diagnose tumor angiogenesis, cancers such as breast cancer, lung cancer, colorectal cancer, renal cancer, pancreatic cancer, or melanoma, ocular diseases or ocular indications, such as diabetic retinopathy, or age related macular degeneration, rheumatoid arthritis, psoriasis, endometriosis, endometrial carcinoma, gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), or menopausal dysfunction.  
25

Such nucleic acid molecules are also useful for the prevention of the diseases and conditions including diabetic retinopathy, macular degeneration, neovascular glaucoma, myopic degeneration, verruca vulgaris, angiomyxoma of tuberous sclerosis, port-wine stains, Sturge Weber syndrome, Kippel-Trenaunay-Weber syndrome, Osler-Weber-Rendu syndrome  
30

and other diseases or conditions that are related to the levels of VEGFR1 or VEGFR2 in a cell or tissue.

In another embodiment, the invention features a composition in a pharmaceutically acceptable carrier or diluent, comprising the nucleic acid molecule of the instant invention.

5        In another embodiment, an enzymatic nucleic acid molecule, antisense nucleic acid molecule, 2-5A antisense chimera, triplex forming oligonucleotide, decoy RNA, dsRNA, siRNA, aptamer, or antisense nucleic acid containing nucleic acid cleaving chemical groups of the invention is adapted for birth control.

10      In one embodiment, an enzymatic nucleic acid molecule of the invention is in a hammerhead, Inozyme, Zinzyme, DNAzyme, Amberzyme, or G-cleaver configuration.

In one embodiment, an enzymatic nucleic acid molecule of the invention comprises between 8 and 100 bases complementary to RNA of VEGFR1 and/or VEGFR2 gene. In another embodiment, an enzymatic nucleic acid molecule of the invention comprises between 14 and 24 bases complementary to RNA of VEGFR1 and/or VEGFR2 gene.

15      In one embodiment, a siRNA molecule of the invention comprises a double stranded RNA wherein one strand of the RNA is complementary to RNA of a VEGFR1 and/or VEGFR2 gene. In another embodiment, a siRNA molecule of the invention comprises a double stranded RNA wherein one strand of the RNA comprises a portion of a sequence of RNA having a VEGFR1 and/or VEGFR2 sequence. In yet another embodiment, a siRNA  
20      molecule of the invention comprises a double stranded RNA wherein both strands of RNA are connected by a non-nucleotide linker. Alternately, a siRNA molecule of the invention comprises a double stranded RNA wherein both strands of RNA are connected by a nucleotide linker, such as a loop or stem loop structure.

25      In one embodiment, a single strand component of a siRNA molecule of the invention is from about 14 to about 50 nucleotides in length. In another embodiment, a single strand component of a siRNA molecule of the invention is about 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, or 28 nucleotides in length. In yet another embodiment, a single strand component of a siRNA molecule of the invention is about 23 nucleotides in length. In one embodiment, a siRNA molecule of the invention is from about 28 to about 56 nucleotides in  
30      length. In another embodiment, a siRNA molecule of the invention is about 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, or 52 nucleotides in length. In yet another embodiment, a siRNA molecule of the invention is about 46 nucleotides in length.

In one embodiment, an enzymatic nucleic acid molecule, antisense nucleic acid molecule, 2-5A antisense chimera, triplex forming oligonucleotide, decoy RNA, dsRNA, siRNA, aptamer, or antisense nucleic acid containing nucleic acid cleaving chemical groups of the invention is chemically synthesized.

5 In another embodiment, an enzymatic nucleic acid molecule, antisense nucleic acid molecule, 2-5A antisense chimera, triplex forming oligonucleotide, decoy RNA, dsRNA, siRNA, aptamer, or antisense nucleic acid containing nucleic acid cleaving chemical groups of the invention comprises at least one 2'-sugar modification.

10 In another embodiment, an enzymatic nucleic acid molecule, antisense nucleic acid molecule, 2-5A antisense chimera, triplex forming oligonucleotide, decoy RNA, dsRNA, siRNA, aptamer, or antisense nucleic acids containing nucleic acid cleaving chemical groups of the invention comprises at least one nucleic acid base modification.

15 In another embodiment, an enzymatic nucleic acid molecule, antisense nucleic acid molecule, 2-5A antisense chimera, triplex forming oligonucleotide, decoy RNA, dsRNA, siRNA, aptamer, or antisense nucleic acid containing nucleic acid cleaving chemical groups of the invention comprises at least one phosphate backbone modification.

In one embodiment, the invention features a mammalian cell, for example a human cell, comprising a nucleic acid molecule of the invention.

20 In another embodiment, the invention features a method of reducing VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 expression or activity in a cell comprising contacting the cell with a nucleic acid molecule of the invention that modulates the expression and/or activity of VEGF and/or VEGFr, under conditions suitable for the reduction.

25 In another embodiment, a method of treatment of a subject having a condition associated with the level of VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 is featured, wherein the method further comprises the use of one or more drug therapies under conditions suitable for the treatment.

30 In one embodiment, the invention features a method for treatment of a subject having tumor angiogenesis, tumor angiogenesis, cancers including but not limited to tumor and cancer types shown under Diagnosis in Table III, ocular diseases or ocular indications such as diabetic retinopathy, or age related macular degeneration, rheumatoid arthritis, psoriasis and/or endometriosis, endometrial carcinoma, gynecologic bleeding disorders, irregular

menstrual cycles, ovulation, premenstrual syndrome (PMS), or menopausal dysfunction, comprising administering to the subject a nucleic acid molecule of the invention that modulates the expression and/or activity of VEGF and/or VEGFr under conditions suitable for the treatment.

5 In another embodiment, the invention features a method for birth control in a subject comprising administering to the subject a nucleic acid molecule of the invention that modulates the expression and/or activity of VEGF and/or VEGFr under conditions suitable for the treatment.

In another embodiment, the invention features a method of cleaving RNA encoded by  
10 a VEGF, VEGFR1 and/or VEGFR2 gene comprising contacting an enzymatic nucleic acid molecule of the invention having endonuclease activity with RNA encoded by a VEGFR1 and/or VEGFR2 gene under conditions suitable for the cleavage, for example, wherein the cleavage is carried out in the presence of a divalent cation, such as  $Mg^{2+}$ .

In one embodiment, a nucleic acid molecule of the invention comprises a cap structure, for example a 3',3'-linked or 5',5'-linked deoxyabasic ribose derivative, wherein  
15 the cap structure is at the 5'-end, or 3'-end, or both the 5'-end and the 3'-end of the enzymatic nucleic acid molecule.

In another embodiment, a nucleic acid molecule of the invention comprises a cap structure, for example a 3',3'-linked or 5',5'-linked deoxyabasic ribose derivative, wherein  
20 the cap structure is at the 5'-end, or 3'-end, or both the 5'-end and the 3'-end of the antisense nucleic acid molecule.

In one embodiment, the invention features an expression vector comprising a nucleic acid sequence encoding at least one nucleic acid molecule of the invention such that the vector allows expression of the nucleic acid molecule.

25 In another embodiment, the invention features a mammalian cell, for example, a human cell comprising an expression vector of the invention.

In yet another embodiment, an expression vector of the invention further comprises a sequence for a nucleic acid molecule complementary to RNA encoded by a VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 gene.

In one embodiment, an expression vector of the invention comprises a nucleic acid sequence encoding two or more nucleic acid molecules of the invention, which can be the same or different.

In another embodiment, the invention features a method for treatment or control of tumor angiogenesis, cancers such as breast cancer, lung cancer, colorectal cancer, renal cancer, pancreatic cancer, or melanoma, or ocular indications such as diabetic retinopathy, or age related macular degeneration, and/or endometriosis, endometrial carcinoma, gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), or menopausal dysfunction, comprising administering to a subject a nucleic acid molecule of the invention that modulates the expression and/or activity of VEGF and/or VEGFr, such as an enzymatic nucleic acid molecule, antisense nucleic acid molecule, 2-5A antisense chimera, triplex forming oligonucleotide, decoy RNA, dsRNA, siRNA, aptamer, or antisense nucleic acid containing nucleic acid cleaving chemical groups of the invention, under conditions suitable for the treatment, including administering to the subject one or more other therapies, for example, 5-fluoro uridine, Leucovorin, Irinotecan (CAMPTOSAR® or CPT-11 or Camptothecin-11 or Campto), Paclitaxel, or Carboplatin. GnRH (gonadotropin releasing hormone) agonists, Lupron Depot (Leuprolide Acetate), Synarel (naferalin acetate), Zolodex (goserelin acetate), Suprefact (buserelin acetate), Danazol, or oral contraceptives including but not limited to Depo-Provera or Provera (medroxyprogesterone acetate), or any other estrogen/progesterone contraceptive.

In one embodiment, the method of treatment features a nucleic acid molecule of the invention, such as an enzymatic nucleic acid or antisense nucleic acid molecule, that comprises at least five ribose residues, at least ten 2'-O-methyl modifications, and a 3'- end modification, such as a 3'-3' inverted abasic moiety. In another embodiment, a nucleic acid molecule of the invention further comprises phosphorothioate linkages on at least three of the 5' terminal nucleotides.

In another embodiment, the invention features a method of administering to a mammal, for example a human, an enzymatic nucleic acid molecule, antisense nucleic acid molecule, 2-5A antisense chimera, triplex forming oligonucleotide, decoy RNA, dsRNA, siRNA, aptamer, or antisense nucleic acid containing nucleic acid cleaving chemical groups of the invention, comprising contacting the mammal with the nucleic acid molecule under conditions suitable for the administration, for example, in the presence of a delivery reagent such as a lipid, cationic lipid, phospholipid, or liposome.

In yet another embodiment, the invention features a method of administering to a mammal an enzymatic nucleic acid molecule, antisense nucleic acid molecule, 2'-5A antisense chimera, triplex forming oligonucleotide, decoy RNA, dsRNA, siRNA, aptamer, or antisense nucleic acid containing nucleic acid cleaving chemical groups of the invention in conjunction  
5 with other therapies, comprising contacting the mammal, for example a human, with the nucleic acid molecule and the other therapy under conditions suitable for the administration.

In another embodiment, other therapies contemplated by the instant invention that can be used in conjunction with the nucleic acid molecules of the instant invention include, but are not limited to, 5-fluoro uridine, Leucovorin, Irinotecan (CAMPTOSAR® or CPT-11 or  
10 Camptothecin-11 or Campto), Paclitaxel, or Carboplatin, GnRH (gonadotropin releasing hormone) agonists, Lupron Depot (Leuprolide Acetate), Synarel (naferalin acetate), Zolodex (goserelin acetate), Suprefact (buserelin acetate), Danazol, or oral contraceptives including but not limited to Depo-Provera or Provera (medroxyprogesterone acetate), or other estrogen/progesterone contraceptive.

15 In one embodiment, the invention features the use of an enzymatic nucleic acid molecule, to down-regulate the expression of VEGFR1 and/or VEGFR2 genes in the treatment or control of tumor angiogenesis, cancers such as breast cancer, lung cancer, colorectal cancer, renal cancer, pancreatic cancer, or melanoma, or ocular indications such as diabetic retinopathy, or age related macular degeneration, and/or endometriosis, endometrial  
20 carcinoma, gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), or menopausal dysfunction. Such enzymatic nucleic acid molecule can be in the hammerhead, NCH, G-cleaver, Amberzyme, Zinzyme, and/or DNAzyme motif.

25 In another embodiment, the invention features the use of an enzymatic nucleic acid molecule to down-regulate the expression of VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 genes, as a method of birth control. Such enzymatic nucleic acid molecule can be in the hammerhead, NCH, G-cleaver, Amberzyme, Zinzyme, and/or DNAzyme motif. In one embodiment, the nucleic acid molecules of the invention have complementarity to the substrate sequences in Tables V and VI. Examples of enzymatic nucleic acid molecules of  
30 the invention are shown in Tables V and VI. Examples of such enzymatic nucleic acid molecules consist essentially of sequences defined in these Tables.

By "inhibit", "down-regulate", or "reduce", it is meant that the expression of the gene, or level of nucleic acids or equivalent nucleic acids encoding one or more proteins or protein subunits, or activity of one or more proteins or protein subunits, such as VEGFR1, VEGFR2

and/or flk-1, is reduced below that observed in the absence of the nucleic acid molecules of the invention. In one embodiment, inhibition, down-regulation or reduction with enzymatic nucleic acid molecule preferably is below that level observed in the presence of an enzymatically inactive or attenuated molecule that is able to bind to the same site on the target 5 nucleic acid, but is unable to cleave that nucleic acid. In another embodiment, inhibition, down-regulation, or reduction with antisense oligonucleotides is preferably below that level observed in the presence of, for example, an oligonucleotide with scrambled sequence or with mismatches. In another embodiment, inhibition, down-regulation, or reduction of VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 with the nucleic acid molecule of the 10 instant invention is greater in the presence of the nucleic acid molecule than in its absence.

By "up-regulate" is meant that the expression of a gene, or level of nucleic acids or equivalent nucleic acids encoding one or more proteins or protein subunits, or activity of one or more proteins or protein subunits, such as VEGFR1 and/or VEGFR2, is greater than that observed in the absence of the nucleic acid molecules of the invention. For example, the 15 expression of a gene, such as VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 gene, can be increased in order to treat, prevent, ameliorate, or modulate a pathological condition caused or exacerbated by an absence or low level of gene expression.

By "modulate" is meant that the expression of a gene, or level of nucleic acids or equivalent nucleic acids encoding one or more proteins or protein subunits, or activity of one 20 or more proteins protein subunit(s) is up-regulated or down-regulated, such that the expression, level, or activity is greater than or less than that observed in the absence of the nucleic acid molecules of the invention.

By "enzymatic nucleic acid molecule" it is meant a nucleic acid molecule which has complementarity in a substrate binding region to a specified gene target, and also has an 25 enzymatic activity which is active to specifically cleave a target nucleic acid. That is, the enzymatic nucleic acid molecule is able to intermolecularly cleave a nucleic acid and thereby inactivate a target nucleic acid molecule. These complementary regions allow sufficient hybridization of the enzymatic nucleic acid molecule to the target nucleic acid and thus permit cleavage. One hundred percent complementarity is preferred, but complementarity as 30 low as 50-75% can also be useful in this invention (see for example Werner and Uhlenbeck, 1995, *Nucleic Acids Research*, 23, 2092-2096; Hammann *et al.*, 1999, *Antisense and Nucleic Acid Drug Dev.*, 9, 25-31). The nucleic acids can be modified at the base, sugar, and/or phosphate groups. The term enzymatic nucleic acid is used interchangeably with phrases such

as ribozymes, catalytic RNA, enzymatic RNA, catalytic DNA, aptazyme or aptamer-binding ribozyme, regulatable ribozyme, catalytic oligonucleotides, nucleozyme, DNAzyme, RNA enzyme, endoribonuclease, endonuclease, minizyme, leadzyme, oligozyme or DNA enzyme. All of these terminologies describe nucleic acid molecules with enzymatic activity. The 5 specific enzymatic nucleic acid molecules described in the instant application are not limiting in the invention and those skilled in the art will recognize that all that is important in an enzymatic nucleic acid molecule of this invention is that it has a specific substrate binding site which is complementary to one or more of the target nucleic acid regions, and that it have nucleotide sequences within or surrounding that substrate binding site which impart a nucleic 10 acid cleaving and/or ligation activity to the molecule (Cech *et al.*, U.S. Patent No. 4,987,071; Cech *et al.*, 1988, 260 *JAMA* 3030).

Several varieties of naturally-occurring enzymatic nucleic acids are known presently. Each can catalyze the hydrolysis of nucleic acid phosphodiester bonds in *trans* (and thus can cleave other nucleic acid molecules) under physiological conditions. Table I summarizes 15 some of the characteristics of these ribozymes. In general, enzymatic nucleic acids act by first binding to a target nucleic acid. Such binding occurs through the target binding portion of a enzymatic nucleic acid which is held in close proximity to an enzymatic portion of the molecule that acts to cleave the target nucleic acid. Thus, the enzymatic nucleic acid first recognizes and then binds a target nucleic acid through complementary base-pairing, and once 20 bound to the correct site, acts enzymatically to cut the target nucleic acid. Strategic cleavage of such a target nucleic acid will destroy its ability to direct synthesis of an encoded protein. After an enzymatic nucleic acid has bound and cleaved its nucleic acid target, it is released from that nucleic acid to search for another target and can repeatedly bind and cleave new targets. Thus, a single ribozyme molecule is able to cleave many molecules of target nucleic 25 acid. In addition, the ribozyme is a highly specific inhibitor of gene expression, with the specificity of inhibition depending not only on the base-pairing mechanism of binding to the target nucleic acid, but also on the mechanism of target nucleic acid cleavage. Single mismatches, or base-substitutions, near the site of cleavage can completely eliminate catalytic activity of a ribozyme.

30 In one embodiment of the inventions described herein, an enzymatic nucleic acid molecule of the invention is formed in a hammerhead or hairpin motif, but can also be formed in the motif of a hepatitis delta virus, group I intron, group II intron or RNase P RNA (in association with an RNA guide sequence), *Neurospora* VS RNA, DNAzymes, NCH cleaving motifs, or G-cleavers. Examples of such hammerhead motifs are described by Dreyfus, 35 *supra*, Rossi *et al.*, 1992, *AIDS Research and Human Retroviruses* 8, 183; of hairpin motifs

by Hampel *et al.*, EP0360257, Hampel and Tritz, 1989 *Biochemistry* 28, 4929, Feldstein *et al.*, 1989, *Gene* 82, 53, Haseloff and Gerlach, 1989, *Gene*, 82, 43, and Hampel *et al.*, 1990 *Nucleic Acids Res.* 18, 299; Chowrira & McSwiggen, US. Patent No. 5,631,359; an examples of a hepatitis delta virus motif is described by Perrotta and Been, 1992 *Biochemistry* 31, 16; 5 examples of RNase P motifs are described by Guerrier-Takada *et al.*, 1983 *Cell* 35, 849; Forster and Altman, 1990, *Science* 249, 783; Li and Altman, 1996, *Nucleic Acids Res.* 24, 835; examples of *Neurospora* VS RNA ribozyme motifs are described by Collins (Saville and Collins, 1990 *Cell* 61, 685-696; Saville and Collins, 1991 *Proc. Natl. Acad. Sci. USA* 88, 8826-8830; Collins and Olive, 1993 *Biochemistry* 32, 2795-2799; Guo and Collins, 1995, 10 *EMBO J.* 14, 363); examples of Group II introns are described by Griffin *et al.*, 1995, *Chem. Biol.* 2, 761; Michels and Pyle, 1995, *Biochemistry* 34, 2965; Pyle *et al.*, International PCT Publication No. WO 96/22689; an example of a Group I intron is described by Cech *et al.*, U.S. Patent 4,987,071; and examples of DNAzymes are described by Usman *et al.*, International PCT Publication No. WO 95/11304; Chartrand *et al.*, 1995, *NAR* 23, 4092; 15 Breaker *et al.*, 1995, *Chem. Bio.* 2, 655; Santoro *et al.*, 1997, *PNAS* 94, 4262, and Beigelman *et al.*, International PCT publication No. WO 99/55857. NCH cleaving motifs are described in Ludwig & Sproat, International PCT Publication No. WO 98/58058; and G-cleavers are described in Kore *et al.*, 1998, *Nucleic Acids Research* 26, 4116-4120 and Eckstein *et al.*, International PCT Publication No. WO 99/16871. Additional motifs such as the Aptazyme 20 (Breaker *et al.*, WO 98/43993), Amberzyme (Beigelman *et al.*, U.S. Serial No. 09/301,511) and Zinzyme (Figure 7) (Beigelman *et al.*, U.S. Serial No. 09/918,728), all included by reference herein including drawings, can also be used in the present invention. These specific motifs or configurations are not limiting in the invention and those skilled in the art will recognize that all that is important in an enzymatic nucleic acid molecule of this invention is 25 that it have a specific substrate binding site which is complementary to one or more of the target gene RNA regions, and that it have nucleotide sequences within or surrounding that substrate binding site which impart a RNA cleaving activity to the molecule (Cech *et al.*, U.S. Patent No. 4,987,071).

By "nucleic acid molecule" as used herein is meant a molecule having nucleotides. The 30 nucleic acid can be single, double, or multiple stranded and can comprise modified or unmodified nucleotides or non-nucleotides or various mixtures and combinations thereof.

By "enzymatic portion" or "catalytic domain" is meant that portion/region of a enzymatic nucleic acid molecule essential for cleavage of a nucleic acid substrate (for example see Figure 6).

By "substrate binding arm" or "substrate binding domain" is meant that portion/region of a enzymatic nucleic acid which is able to interact, for example via complementarity (*i.e.*, able to base-pair with), with a portion of its substrate. Preferably, such complementarity is 100%, but can be less if desired. For example, as few as 10 bases out of 14 can be base-paired 5 (see for example Werner and Uhlenbeck, 1995, *Nucleic Acids Research*, 23, 2092-2096; Hammann *et al.*, 1999, *Antisense and Nucleic Acid Drug Dev.*, 9, 25-31). Examples of such arms are shown generally in Figures 6-8. That is, these arms contain sequences within a enzymatic nucleic acid which are intended to bring enzymatic nucleic acid and target nucleic acid together through complementary base-pairing interactions. An enzymatic nucleic acid of 10 the invention can have binding arms that are contiguous or non-contiguous and can be of varying lengths. The length of the binding arm(s) are preferably greater than or equal to four nucleotides and of sufficient length to stably interact with the target nucleic acid; preferably 12-100 nucleotides; more preferably 14-24 nucleotides long (see for example Werner and Uhlenbeck, *supra*; Hamman *et al.*, *supra*; Hampel *et al.*, EP0360257; Berzal-Herranz *et al.*, 15 1993, *EMBO J.*, 12, 2567-73) or between 8 and 14 nucleotides long. If two binding arms are chosen, the design is such that the length of the binding arms are symmetrical (*i.e.*, each of the binding arms is of the same length; *e.g.*, four and four, five and five nucleotides, or six and six nucleotides, or seven and seven nucleotides long) or asymmetrical (*i.e.*, the binding arms are of different length; *e.g.*, three and five, six and three nucleotides; three and six 20 nucleotides long; four and five nucleotides long; four and six nucleotides long; four and seven nucleotides long; and the like).

By "Inozyme" or "NCH" motif or configuration is meant, an enzymatic nucleic acid molecule comprising a motif as is generally described as NCH Rz in Figure 6 and in Ludwig *et al.*, International PCT Publication No. WO 98/58058 and US Patent Application Serial No. 25 08/878,640. Inozymes possess endonuclease activity to cleave nucleic acid substrates having a cleavage triplet NCH/, where N is a nucleotide, C is cytidine and H is adenosine, uridine or cytidine, and "/" represents the cleavage site. H is used interchangeably with X. Inozymes can also possess endonuclease activity to cleave nucleic acid substrates having a cleavage triplet NCN/, where N is a nucleotide, C is cytidine, and "/" represents the cleavage site. "T" 30 in Figure 6 represents an Inosine nucleotide, preferably a ribo-Inosine or xylo-Inosine nucleoside.

By "G-cleaver" motif or configuration is meant, an enzymatic nucleic acid molecule comprising a motif as is generally described as G-cleaver Rz in Figure 6 and in Eckstein *et al.*, US 6,127,173. G-cleavers possess endonuclease activity to cleave nucleic acid substrates 35 having a cleavage triplet NYN/, where N is a nucleotide, Y is uridine or cytidine and "/"

represents the cleavage site. G-cleavers can be chemically modified as is generally shown in Figure 6.

By "amberzyme" motif or configuration is meant, an enzymatic nucleic acid molecule comprising a motif as is generally described in Beigelman *et al.*, International PCT publication No. WO 99/55857 and US Patent Application Serial No. 09/476,387. 5 Amberzymes possess endonuclease activity to cleave nucleic acid substrates having a cleavage triplet NG/N, where N is a nucleotide, G is guanosine, and "/" represents the cleavage site. Amberzymes can be chemically modified to increase nuclease stability through substitutions using modified nucleotides. In addition, differing nucleoside and/or non-10 nucleoside linkers can be used to substitute the 5'-gaaa-3' loops shown in the figure. Amberzymes represent a non-limiting example of an enzymatic nucleic acid molecule that does not require a ribonucleotide (2'-OH) group within its own nucleic acid sequence for activity.

By "zinzyme" motif or configuration is meant, an enzymatic nucleic acid molecule comprising a motif as is generally described in Figure 7 and in Beigelman *et al.*, International PCT publication No. WO 99/55857 and US Patent Application Serial No. 09/918,728. 15 Zinzymes possess endonuclease activity to cleave nucleic acid substrates having a cleavage triplet including but not limited to YG/Y, where Y is uridine or cytidine, and G is guanosine and "/" represents the cleavage site. Zinzymes can be chemically modified to increase nuclease stability through substitutions as are generally shown in Figure 7, including substituting 2'-O-methyl guanosine nucleotides for guanosine nucleotides. In addition, differing nucleotide and/or non-nucleotide linkers can be used to substitute the 5'-gaaa-2' 20 loop shown in the figure. Zinzymes represent a non-limiting example of an enzymatic nucleic acid molecule that does not require a ribonucleotide (2'-OH) group within its own nucleic acid sequence for activity.

By 'DNAzyme' is meant, an enzymatic nucleic acid molecule that does not require the presence of a 2'-OH group within its own nucleic acid sequence for activity. In particular embodiments the enzymatic nucleic acid molecule can have an attached linker or linkers or other attached or associated groups, moieties, or chains containing one or more nucleotides 30 with 2'-OH groups. DNAzymes can be synthesized chemically or expressed endogenously *in vivo*, by means of a single stranded DNA vector or equivalent thereof. An example of a DNAzyme is shown in Figure 8 and is generally reviewed in Usman *et al.*, US patent No., 6,159,714; Chartrand *et al.*, 1995, *NAR* 23, 4092; Breaker *et al.*, 1995, *Chem. Bio.* 2, 655; Santoro *et al.*, 1997, *PNAS* 94, 4262; Breaker, 1999, *Nature Biotechnology*, 17, 422-423; and

Santoro *et. al.*, 2000, *J. Am. Chem. Soc.*, 122, 2433-39. The "10-23" DNAzyme motif is one particular type of DNAzyme that was evolved using *in vitro* selection, see Santoro *et al.*, *supra* and as generally described in Joyce *et al.*, US 5,807,718. Additional DNAzyme motifs can be selected for using techniques similar to those described in these references, and hence, 5 are within the scope of the present invention.

By "sufficient length" is meant a nucleic acid molecule of the invention is long enough to provide the intended function under the expected condition. For example, a nucleic acid molecule of the invention needs to be of "sufficient length" to provide stable interaction with a target nucleic acid molecule under the expected binding conditions and environment. In 10 another non-limiting example, for the binding arms of an enzymatic nucleic acid, "sufficient length" means that the binding arm sequence is long enough to provide stable binding to a target site under the expected reaction conditions and environment. The binding arms are not so long as to prevent useful turnover of the nucleic acid molecule.

By "stably interact" is meant interaction of an oligonucleotides with target nucleic acid 15 (*e.g.*, by forming hydrogen bonds with complementary nucleotides in the target under physiological conditions) that is sufficient to the intended purpose (*e.g.*, cleavage of target nucleic acid by an enzyme).

By "equivalent" RNA to VEGF, VEGFR1 and/or VEGFR2 is meant to include nucleic acid molecules having homology (partial or complete) to a nucleic acid encoding VEGF, 20 VEGFR1 and/or VEGFR2 proteins or encoding proteins with similar function as VEGF, VEGFR1 and/or VEGFR2 proteins in various organisms, including human, rodent, primate, rabbit, pig, protozoans, fungi, plants, and other microorganisms and parasites. The equivalent nucleic acid sequence also includes, in addition to the coding region, regions such as 5'-untranslated region, 3'-untranslated region, introns, intron-exon junction and the like.

25 By "homology" is meant the nucleotide sequence of two or more nucleic acid molecules is partially or completely identical.

By "antisense nucleic acid", it is meant a non-enzymatic nucleic acid molecule that binds to target nucleic acid by means of RNA-RNA or RNA-DNA or RNA-PNA (protein nucleic acid; Egholm *et al.*, 1993 *Nature* 365, 566) interactions and alters the activity of the 30 target nucleic acid (for a review, see Stein and Cheng, 1993 *Science* 261, 1004 and Woolf *et al.*, US patent No. 5,849,902). Typically, antisense molecules are complementary to a target sequence along a single contiguous sequence of the antisense molecule. However, in certain embodiments, an antisense molecule can bind to substrate such that the substrate molecule

forms a loop, and/or an antisense molecule can bind such that the antisense molecule forms a loop. Thus, an antisense molecule can be complementary to two (or even more) non-contiguous substrate sequences or two (or even more) non-contiguous sequence portions of an antisense molecule can be complementary to a target sequence or both. For a review of 5 current antisense strategies, see Schmajuk *et al.*, 1999, *J. Biol. Chem.*, 274, 21783-21789, Delihas *et al.*, 1997, *Nature*, 385, 751-753, Stein *et al.*, 1997, *Antisense N. A. Drug Dev.*, 7, 151, Crooke, 2000, *Methods Enzymol.*, 313, 3-45; Crooke, 1998, *Biotech. Genet. Eng. Rev.*, 15, 121-157, Crooke, 1997, *Ad. Pharmacol.*, 40, 1-49. In addition, antisense DNA can be used to target nucleic acid by means of DNA-RNA interactions, thereby activating RNase H, 10 which digests the target nucleic acid in the duplex. The antisense oligonucleotides can comprise one or more RNase H activating region, which is capable of activating RNase H cleavage of a target nucleic acid. Antisense DNA can be synthesized chemically or expressed via the use of a single stranded DNA expression vector or equivalent thereof.

By "RNase H activating region" is meant a region (generally greater than or equal to 4- 15 25 nucleotides in length, preferably from 5-11 nucleotides in length) of a nucleic acid molecule capable of binding to a target nucleic acid to form a non-covalent complex that is recognized by cellular RNase H enzyme (see for example Arrow *et al.*, US 5,849,902; Arrow *et al.*, US 5,989,912). The RNase H enzyme binds to a nucleic acid molecule-target nucleic acid complex and cleaves the target nucleic acid sequence. The RNase H activating region 20 comprises, for example, phosphodiester, phosphorothioate (preferably at least four of the nucleotides are phosphorothioate substitutions; more specifically, 4-11 of the nucleotides are phosphorothioate substitutions); phosphorodithioate, 5'-thiophosphate, or methylphosphonate backbone chemistry or a combination thereof. In addition to one or more backbone chemistries described above, the RNase H activating region can also comprise a variety of 25 sugar chemistries. For example, the RNase H activating region can comprise deoxyribose, arabino, fluoroarabino or a combination thereof, nucleotide sugar chemistry. Those skilled in the art will recognize that the foregoing are non-limiting examples and that any combination of phosphate, sugar and base chemistry of a nucleic acid that supports the activity of RNase H enzyme is within the scope of the definition of the RNase H activating region and the instant 30 invention.

By "2-5A antisense chimera" is meant an antisense oligonucleotide containing a 5'- phosphorylated 2'-5'-linked adenylate residue. These chimeras bind to target nucleic acid in a sequence-specific manner and activate a cellular 2-5A-dependent ribonuclease which, in turn, cleaves the target nucleic acid (Torrence *et al.*, 1993 *Proc. Natl. Acad. Sci. USA* 90, 1300;

Silverman *et al.*, 2000, *Methods Enzymol.*, 313, 522-533; Player and Torrence, 1998, *Pharmacol. Ther.*, 78, 55-113).

By "triplex forming oligonucleotides" is meant an oligonucleotide that can bind to a double-stranded polynucleotide, such as DNA, in a sequence-specific manner to form a triple-strand helix. Formation of such triple helix structure has been shown to inhibit transcription of the targeted gene (Duval-Valentin *et al.*, 1992 *Proc. Natl. Acad. Sci. USA* 89, 504; Fox, 2000, *Curr. Med. Chem.*, 7, 17-37; Praseuth *et. al.*, 2000, *Biochim. Biophys. Acta*, 1489, 181-206).

By "gene" it is meant a nucleic acid that encodes an RNA, for example, nucleic acid sequences including but not limited to structural genes encoding a polypeptide.

The term "complementarity" as used herein refers to the ability of a nucleic acid to form hydrogen bond(s) with another nucleic acid sequence by either traditional Watson-Crick or other non-traditional types. In reference to nucleic molecules of the present invention, the binding free energy for a nucleic acid molecule with its target or complementary sequence is sufficient to allow the relevant function of the nucleic acid to proceed, e.g., enzymatic nucleic acid cleavage, antisense or triple helix inhibition. Determination of binding free energies for nucleic acid molecules is well known in the art (see, e.g., Turner *et al.*, 1987, *CSH Symp. Quant. Biol.* LII pp.123-133; Frier *et al.*, 1986, *Proc. Nat. Acad. Sci. USA* 83:9373-9377; Turner *et al.*, 1987, *J. Am. Chem. Soc.* 109:3783-3785). A percent complementarity indicates the percentage of contiguous residues in a nucleic acid molecule which can form hydrogen bonds (e.g., Watson-Crick base pairing) with a second nucleic acid sequence (e.g., 5, 6, 7, 8, 9, 10 out of 10 being 50%, 60%, 70%, 80%, 90%, and 100% complementary). "Perfectly complementary" means that all the contiguous residues of a nucleic acid sequence will hydrogen bond with the same number of contiguous residues in a second nucleic acid sequence.

By "RNA" is meant a molecule comprising at least one ribonucleotide residue. By "ribonucleotide" or "2'-OH" is meant a nucleotide with a hydroxyl group at the 2' position of a β-D-ribo-furanose moiety.

By "nucleic acid decoy molecule", or "decoy" as used herein is meant a nucleic acid molecule that mimics the natural binding domain for a ligand. The decoy therefore competes with the natural binding target for the binding of a specific ligand. For example, it has been shown that over-expression of HIV trans-activation response (TAR) RNA can act as a

"decoy" and efficiently binds HIV tat protein, thereby preventing it from binding to TAR sequences encoded in the HIV RNA (Sullenger et al., 1990, *Cell*, 63, 601-608).

By "aptamer" or "nucleic acid aptamer" as used herein is meant a nucleic acid molecule that binds specifically to a target molecule wherein the nucleic acid molecule has sequence 5 that is distinct from sequence recognized by the target molecule in its natural setting. Alternately, an aptamer can be a nucleic acid molecule that binds to a target molecule where the target molecule does not naturally bind to a nucleic acid. The target molecule can be any molecule of interest. For example, the aptamer can be used to bind to a ligand binding domain of a protein, thereby preventing interaction of the naturally occurring ligand with the protein. 10 Similarly, the nucleic acid molecules of the instant invention can bind to VEGFR1 or VEGFR2 receptors to block activity of the receptor. This is a non-limiting example and those in the art will recognize that other embodiments can be readily generated using techniques generally known in the art, see for example Gold et al., US 5,475,096 and 5,270,163; Gold et al., 1995, *Annu. Rev. Biochem.*, 64, 763; Brody and Gold, 2000, *J. Biotechnol.*, 74, 5; Sun, 15 2000, *Curr. Opin. Mol. Ther.*, 2, 100; Kusser, 2000, *J. Biotechnol.*, 74, 27; Hermann and Patel, 2000, *Science*, 287, 820; and Jayasena, 1999, *Clinical Chemistry*, 45, 1628.

The term "double stranded RNA" or "dsRNA" as used herein refers to a double stranded RNA molecule capable of RNA interference "RNAi", including short interfering RNA "siRNA" see for example Bass, 2001, *Nature*, 411, 428-429; Elbashir et al., 2001, 20 *Nature*, 411, 494-498; and Kreutzer et al., International PCT Publication No. WO 00/44895; Zernicka-Goetz et al., International PCT Publication No. WO 01/36646; Fire, International PCT Publication No. WO 99/32619; Plaetinck et al., International PCT Publication No. WO 00/01846; Mello and Fire, International PCT Publication No. WO 01/29058; Deschamps-Depaillette, International PCT Publication No. WO 99/07409; and Li et al., International PCT 25 Publication No. WO 00/44914.

By "nucleic acid sensor molecule" or "allozyme" as used herein is meant a nucleic acid molecule comprising an enzymatic domain and a sensor domain, where the enzymatic nucleic acid domain's ability to catalyze a chemical reaction is dependent on the interaction with a target signaling molecule, such as a nucleic acid, polynucleotide, oligonucleotide, 30 peptide, polypeptide, or protein, for example VEGF, VEGFR1 and/or VEGFR2. The introduction of chemical modifications, additional functional groups, and/or linkers, to the nucleic acid sensor molecule can provide enhanced catalytic activity of the nucleic acid sensor molecule, increased binding affinity of the sensor domain to a target nucleic acid, and/or improved nuclease/chemical stability of the nucleic acid sensor molecule, and are

- hence within the scope of the present invention (see for example Usman *et al.*, US Patent Application No. 09/877,526, George *et al.*, US Patent Nos. 5,834,186 and 5,741,679, Shih *et al.*, US Patent No. 5,589,332, Nathan *et al.*, US Patent No 5,871,914, Nathan and Ellington, International PCT publication No. WO 00/24931, Breaker *et al.*, International PCT Publication Nos. WO 00/26226 and 98/27104, and Sullenger *et al.*, US Patent Application Serial No. 09/205,520).

By "sensor component" or "sensor domain" of the nucleic acid sensor molecule as used herein is meant, a nucleic acid sequence (e.g., RNA or DNA or analogs thereof) which interacts with a target signaling molecule, for example a nucleic acid sequence in one or more regions of a target nucleic acid molecule or more than one target nucleic acid molecule, and which interaction causes the enzymatic nucleic acid component of the nucleic acid sensor molecule to either catalyze a reaction or stop catalyzing a reaction. In the presence of target signaling molecule of the invention, such as VEGF, VEGFR1 and/or VEGFR2, the ability of the sensor component, for example, to modulate the catalytic activity of the nucleic acid sensor molecule, is inhibited or diminished. The sensor component can comprise recognition properties relating to chemical or physical signals capable of modulating the nucleic acid sensor molecule via chemical or physical changes to the structure of the nucleic acid sensor molecule. The sensor component can be derived from a naturally occurring nucleic acid binding sequence, for example, RNAs that bind to other nucleic acid sequences *in vivo*.

Alternately, the sensor component can be derived from a nucleic acid molecule (aptamer) which is evolved to bind to a nucleic acid sequence within a target nucleic acid molecule (see for example Gold *et al.*, US 5,475,096 and 5,270,163). The sensor component can be covalently linked to the nucleic acid sensor molecule, or can be non-covalently associated. A person skilled in the art will recognize that all that is required is that the sensor component is able to selectively inhibit the activity of the nucleic acid sensor molecule to catalyze a reaction.

By "target molecule" or "target signaling molecule" is meant a molecule capable of interacting with a nucleic acid sensor molecule, specifically a sensor domain of a nucleic acid sensor molecule, in a manner that causes the nucleic acid sensor molecule to be active or inactive. The interaction of the signaling agent with a nucleic acid sensor molecule can result in modification of the enzymatic nucleic acid component of the nucleic acid sensor molecule via chemical, physical, topological, or conformational changes to the structure of the molecule, such that the activity of the enzymatic nucleic acid component of the nucleic acid sensor molecule is modulated, for example is activated or deactivated. Signaling agents can comprise target signaling molecules such as macromolecules, ligands, small molecules,

metals and ions, nucleic acid molecules including but not limited to RNA and DNA or analogs thereof, proteins, peptides, antibodies, polysaccharides, lipids, sugars, microbial or cellular metabolites, pharmaceuticals, and organic and inorganic molecules in a purified or unpurified form, for example VEGF, VEGFR1 and/or VEGFR2.

- 5       The term "triplex forming oligonucleotides" as used herein refers to an oligonucleotide that can bind to a double-stranded DNA in a sequence-specific manner to form a triple-strand helix. Formation of such a triple helix structure has been shown to inhibit transcription of a targeted gene (Duval-Valentin *et al.*, 1992 *Proc. Natl. Acad. Sci. USA* 89, 504; Fox, 2000, *Curr. Med. Chem.*, 7, 17-37; Praseuth *et. al.*, 2000, *Biochim. Biophys. Acta*, 1489, 181-206).
- 10      The nucleic acid molecules that modulate the expression of VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 specific nucleic acids, represent a novel therapeutic approach to treat or control a variety of angiogenesis related disorders and conditions, including but not limited to tumor angiogenesis, cancers such as breast cancer, lung cancer, colorectal cancer, renal cancer, pancreatic cancer, or melanoma, or ocular indications such as diabetic 15     retinopathy, or age related macular degeneration, and/or endometriosis, endometrial carcinoma, gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), and/or menopausal dysfunction. The nucleic acid molecules that modulate the expression of VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 specific nucleic acids also represent a novel approach to control ovulation or embryonic 20     implantation and therefore provide a novel means of birth control.

In one embodiment of the present invention, a nucleic acid molecule of the instant invention can be between 12 and 100 nucleotides in length. An exemplary enzymatic nucleic acid molecule of the invention is shown as Formula I and/or Formula II. For example, enzymatic nucleic acid molecules of the invention are preferably between 15 and 50 25     nucleotides in length, more preferably between 25 and 40 nucleotides in length, e.g., 34, 36, or 38 nucleotides in length (for example see Jarvis *et al.*, 1996, *J. Biol. Chem.*, 271, 29107-29112). Exemplary DNAzymes of the invention are preferably between 15 and 40 nucleotides in length, more preferably between 25 and 35 nucleotides in length, e.g., 29, 30, 31, or 32 nucleotides in length (see for example Santoro *et al.*, 1998, *Biochemistry*, 37, 13330-13342; 30     Chartrand *et al.*, 1995, *Nucleic Acids Research*, 23, 4092-4096). Exemplary antisense molecules of the invention are preferably between 15 and 75 nucleotides in length, more preferably between 20 and 35 nucleotides in length, e.g., 25, 26, 27, or 28 nucleotides in length (see for example Woolf *et al.*, 1992, *PNAS*, 89, 7305-7309; Milner *et al.*, 1997, *Nature Biotechnology*, 15, 537-541). Exemplary triplex forming oligonucleotide molecules 35     of the invention are preferably between 10 and 40 nucleotides in length, more preferably

between 12 and 25 nucleotides in length, e.g., 18, 19, 20, or 21 nucleotides in length (see for example Maher *et al.*, 1990, *Biochemistry*, 29, 8820-8826; Strobel and Dervan, 1990, *Science*, 249, 73-75). Those skilled in the art will recognize that all that is required is that the nucleic acid molecule be of length and conformation sufficient and suitable for the nucleic  
5 acid molecule to catalyze a reaction contemplated herein. The length of the nucleic acid molecules of the instant invention are not limiting within the general limits stated.

In a preferred embodiment, a nucleic acid molecule that modulates, for example, down-regulates, VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 replication or expression comprises between 8 and 100 bases complementary to a nucleic acid molecule of  
10 VEGFR1 and/or VEGFR2. More preferably, a nucleic acid molecule that modulates VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 replication or expression comprises between 14 and 24 bases complementary to a nucleic acid molecule of VEGFR1 and/or VEGFR2.

The invention provides a method for producing a class of nucleic acid-based gene  
15 modulating agents which exhibit a high degree of specificity for the nucleic acid of a desired target. For example, a nucleic acid molecule of the invention is preferably targeted to a highly conserved sequence region of target nucleic acids encoding VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 (specifically VEGF, VEGFR1 and/or VEGFR2 genes) such that specific treatment of a disease or condition can be provided with either one or several nucleic  
20 acid molecules of the invention. Such nucleic acid molecules can be delivered exogenously to specific tissue or cellular targets as required. Alternatively, the nucleic acid molecules can be expressed from DNA and/or RNA vectors that are delivered to specific cells.

As used in herein "cell" is used in its usual biological sense, and does not refer to an entire multicellular organism. The cell can, for example, be *in vitro*, e.g., in cell culture, or  
25 present in a multicellular organism, including, e.g., birds, plants and mammals such as humans, cows, sheep, apes, monkeys, swine, dogs, and cats. The cell may be prokaryotic (e.g., bacterial cell) or eukaryotic (e.g., mammalian or plant cell).

By "VEGFR1 and/or VEGFR2 proteins" is meant, protein receptor or a mutant protein derivative thereof, having vascular endothelial growth factor receptor activity, for example,  
30 having the ability to bind vascular endothelial growth factor and/or having tyrosine kinase activity.

By "highly conserved sequence region" is meant, a nucleotide sequence of one or more regions in a target gene does not vary significantly from one generation to the other or from one biological system to the other.

"Angiogenesis" refers to formation of new blood vessels which is an essential process  
5 in reproduction, development and wound repair. "Tumor angiogenesis" refers to the induction of the growth of blood vessels from surrounding tissue into a solid tumor. Tumor growth and tumor metastasis are dependent on angiogenesis (for a review see Folkman, 1985 *supra*; Folkman 1990 *J. Natl. Cancer Inst.*, 82, 4; Folkman and Shing, 1992 *J. Biol. Chem.* 267, 10931).

10 Angiogenesis plays an important role in other diseases such as arthritis wherein new blood vessels have been shown to invade the joints and degrade cartilage (Folkman and Shing, *supra*).

"Retinopathy" refers to inflammation of the retina and/or degenerative condition of the retina which may lead to occlusion of the retina and eventual blindness. In "diabetic 15 retinopathy" angiogenesis causes the capillaries in the retina to invade the vitreous resulting in bleeding and blindness which is also seen in neonatal retinopathy (for a review see Folkman, 1985 *supra*; Folkman 1990 *supra*; Folkman and Shing, 1992 *supra*).

Nucleic acid-based inhibitors of VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2, expression are useful for the prevention, treatment, and/or control of angiogenesis 20 related disorders and conditions, including but not limited to, tumor angiogenesis, cancers such as breast cancer, lung cancer, colorectal cancer, renal cancer, pancreatic cancer, or melanoma, or ocular indications such as diabetic retinopathy, or age related macular degeneration, and/or endometriosis, endometrial carcinoma, gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), menopausal 25 dysfunction, and other diseases or conditions that are related to or will respond to the levels of VEGF, VEGFR1 and/or VEGFR2 in a cell or tissue, alone or in combination with other therapies. The reduction of VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 expression (specifically VEGF, VEGFR1 and/or VEGFR2 gene RNA levels) and thus reduction in the level of the respective protein relieves, to some degree, the symptoms of the 30 disease or condition. Nucleic acid-based inhibitors of VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 expression are also useful as birth control agents, for example by inhibition of ovulation or embryonic uterine implantation.

The nucleic acid molecules of the invention can be added directly, or can be complexed with cationic lipids, packaged within liposomes, or otherwise delivered to target cells or tissues. The nucleic acid complexes can be locally administered to relevant tissues ex vivo, or in vivo through injection or infusion pump, with or without their incorporation in 5 biopolymers. In preferred embodiments, the nucleic acid inhibitors comprise sequences, which are complementary to polynucleotides, for example DNA and RNA, having VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 sequence.

Triplex molecules of the invention can be provided targeted to DNA target regions, and containing the DNA equivalent of a target sequence or a sequence complementary to the 10 specified target (substrate) sequence. Antisense molecules typically are complementary to a target sequence along a single contiguous sequence of the antisense molecule. However, in certain embodiments, an antisense molecule can bind to substrate such that the substrate molecule forms a loop, and/or an antisense molecule can bind such that the antisense molecule forms a loop. Thus, the antisense molecule can be complementary to two (or even 15 more) non-contiguous substrate sequences or two (or even more) non-contiguous sequence portions of an antisense molecule can be complementary to a target sequence or both.

By "consists essentially of" is meant that the active nucleic acid molecule of the invention, for example, an enzymatic nucleic acid molecule, contains an enzymatic center or core equivalent to those in the examples, and binding arms able to bind nucleic acid such that 20 cleavage at the target site occurs. Other sequences can be present which do not interfere with such cleavage. Thus, a core region can, for example, include one or more loop, stem-loop structure, or linker which does not prevent enzymatic activity. Thus, a particular region of a nucleic acid molecule of the invention can be such a loop, stem-loop, nucleotide linker, and/or non-nucleotide linker and can be represented generally as sequence "X". Thus, a core 25 region may, for example, include one or more loop or stem-loop structures which do not prevent enzymatic activity. For example, a core sequence for a hammerhead enzymatic nucleic acid can comprise a conserved sequence, such as 5'-CUGAUGAG-3' and 5'-CGAA-3' connected by "X", where X is 5'-GCCGUUAGGC-3' (SEQ ID NO 5979), or any other Stem II region known in the art, or a nucleotide and/or non-nucleotide linker. Similarly, for 30 other nucleic acid molecules of the instant invention, such as Inozyme, G-cleaver, amberzyme, zinzyme, DNAzyme, antisense, 2-5A antisense, triplex forming nucleic acid, aptamers, decoy nucleic acids, dsRNA or siRNA, other sequences or non-nucleotide linkers can be present that do not interfere with the function of the nucleic acid molecule.

Sequence X can be a linker of  $\geq$  2 nucleotides in length, preferably 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 26, 30, where the nucleotides can preferably be internally base-paired to form a stem or preferably  $\geq$  2 base pairs. Alternatively or in addition, sequence X can be a non-nucleotide linker. In yet another embodiment, the nucleotide linker X can be a nucleic acid aptamer, such 5 as an ATP aptamer, HIV Rev aptamer (RRE), HIV Tat aptamer (TAR) and others (for a review see Gold *et al.*, 1995, *Annu. Rev. Biochem.*, 64, 763; and Szostak & Ellington, 1993, in *The RNA World*, ed. Gesteland and Atkins, pp. 511, CSH Laboratory Press). A nucleic acid aptamer includes a nucleic acid sequence capable of interacting with a ligand. The ligand can be any natural or a synthetic molecule, including but not limited to a resin, metabolites, 10 nucleosides, nucleotides, drugs, toxins, transition state analogs, peptides, lipids, proteins, amino acids, nucleic acid molecules, hormones, carbohydrates, receptors, cells, viruses, bacteria and others.

In yet another embodiment, the non-nucleotide linker X is as defined herein. The term "non-nucleotide" as used herein include either abasic nucleotide, polyether, polyamine, 15 polyamide, peptide, carbohydrate, lipid, or polyhydrocarbon compounds. Specific examples include those described by Seela and Kaiser, *Nucleic Acids Res.* 1990, 18:6353 and *Nucleic Acids Res.* 1987, 15:3113; Cload and Schepartz, *J. Am. Chem. Soc.* 1991, 113:6324; Richardson and Schepartz, *J. Am. Chem. Soc.* 1991, 113:5109; Ma *et al.*, *Nucleic Acids Res.* 1993, 21:2585 and *Biochemistry* 1993, 32:1751; Durand *et al.*, *Nucleic Acids Res.* 1990, 18:6353; McCurdy *et al.*, *Nucleosides & Nucleotides* 1991, 10:287; Jschke *et al.*, *Tetrahedron Lett.* 1993, 34:301; Ono *et al.*, *Biochemistry* 1991, 30:9914; Arnold *et al.*, International Publication No. WO 89/02439; Usman *et al.*, International Publication No. WO 95/06731; Dudycz *et al.*, International Publication No. WO 95/11910 and Ferentz and Verdine, *J. Am. Chem. Soc.* 1991, 113:4000, all hereby incorporated by reference herein.

25 A "non-nucleotide" further means any group or compound which can be incorporated into a nucleic acid chain in the place of one or more nucleotide units, including either sugar and/or phosphate substitutions, and allows the remaining bases to exhibit their enzymatic activity. The group or compound can be abasic in that it does not contain a commonly recognized nucleotide base, such as adenosine, guanine, cytosine, uracil or thymine. Thus, in 30 one embodiment, the invention features an enzymatic nucleic acid molecule having one or more non-nucleotide moieties, and having enzymatic activity to cleave an RNA or DNA molecule.

In another aspect of the invention, nucleic acid molecules that interact with target nucleic acid molecules and down-regulate VEGF and/or VEGFr, such as VEGFR1 and/or

VEGFR2 (specifically VEGF, VEGFR1 and/or VEGFR2 gene) activity are expressed from transcription units inserted into DNA or RNA vectors. The recombinant vectors are preferably DNA plasmids or viral vectors. Enzymatic nucleic acid molecule or antisense expressing viral vectors can be constructed based on, but not limited to, adeno-associated virus, retrovirus, adenovirus, or alphavirus. The recombinant vectors capable of expressing the enzymatic nucleic acid molecules or antisense are delivered as described above, and persist in target cells. Alternatively, viral vectors can be used that provide for transient expression of enzymatic nucleic acid molecules or antisense. Such vectors can be repeatedly administered as necessary. Once expressed, the enzymatic nucleic acid molecules or antisense bind to the target nucleic acid and down-regulate its function or expression. Delivery of enzymatic nucleic acid molecule or antisense expressing vectors can be systemic, such as by intravenous or intramuscular administration, by administration to target cells explanted from the patient followed by reintroduction into the patient, or by any other means that would allow for introduction into the desired target cell. Antisense DNA can be expressed via the use of a single stranded DNA intracellular expression vector.

By "vectors" is meant any nucleic acid- and/or viral-based technique used to deliver a desired nucleic acid.

By "subject" or "patient" is meant an organism, which is a donor or recipient of explanted cells, or the cells themselves. "Subject" or "Patient" also refers to an organism to which the nucleic acid molecules of the invention can be administered. Preferably, a subject or patient is a mammal or mammalian cells. More preferably, a subject or patient is a human or human cells.

By "enhanced enzymatic activity" is meant to include activity measured in cells and/or *in vivo* where the activity is a reflection of both the catalytic activity and the stability of the nucleic acid molecules of the invention. In this invention, the product of these properties can be increased *in vivo* compared to an all RNA enzymatic nucleic acid or all DNA enzyme. In some cases, the activity or stability of the nucleic acid molecule can be decreased (i.e., less than ten-fold), but the overall activity of the nucleic acid molecule is enhanced, *in vivo*.

The nucleic acid molecules of the instant invention, individually, or in combination or in conjunction with other drugs, can be used to treat diseases or conditions discussed above. For example, to treat a disease or condition associated with the levels of VEGFR1 and/or VEGFR2, the patient can be treated, or other appropriate cells can be treated, as is evident to those skilled in the art, individually or in combination with one or more drugs under conditions suitable for the treatment.

In a further embodiment, the described molecules of the invention can be used in combination with other known treatments to treat conditions or diseases discussed above. For example, the described molecules can be used in combination with one or more known therapeutic agents to treat angiogenesis related disorders and conditions, including but not limited to tumor angiogenesis, cancers such as breast cancer, lung cancer, colorectal cancer, renal cancer, pancreatic cancer, or melanoma, or ocular indications such as diabetic retinopathy, or age related macular degeneration, and/or endometriosis, birth control, endometrial tumors, gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), menopausal dysfunction, endometrial carcinoma, and/or other diseases or conditions which respond to the modulation of VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 expression.

Other features and advantages of the invention will be apparent from the following description of the preferred embodiments thereof, and from the claims.

15

Brief Description of the Drawings

**Figure 1** shows a secondary structure model of ANGIOZYME™ ribozyme bound to its RNA target.

**Figure 2** shows a time course of inhibition of primary tumor growth following systemic administration of ANGIOZYME™ in the LLC mouse model.

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**Figure 3** shows inhibition of primary tumor growth following systemic administration of ANGIOZYME™ according to a certain dosing regimen in the LLC mouse model.

**Figure 4** shows a dose-dependent inhibition of tumor metastases following systemic administration of ANGIOZYME™ in a mouse colorectal model.

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**Figure 5** is a graph showing the plasma concentration profile of ANGIOZYME™ after a single subcutaneous (SC) dose of 10, 30, 100 or 300 mg/m<sup>2</sup>.

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**Figure 6** shows examples of chemically stabilized ribozyme motifs. HH Rz, represents hammerhead ribozyme motif (Usman *et al.*, 1996, *Curr. Op. Struct. Bio.*, 1, 527); NCH Rz represents the NCH ribozyme motif (Ludwig *et al.*, International PCT Publication No. WO 98/58058 and US Patent Application Serial No. 08/878,640); G-Cleaver, represents G-cleaver ribozyme motif (Kore *et al.*, 1998, *Nucleic Acids Research* 26, 4116-4120, Eckstein *et*

*al.*, US 6,127,173). N or n, represent independently a nucleotide which can be same or different and have complementarity to each other; rI, represents ribo-Inosine nucleotide; arrow indicates the site of cleavage within the target. Position 4 of the HH Rz and the NCH Rz is shown as having 2'-C-allyl modification, but those skilled in the art will recognize that 5 this position can be modified with other modifications well known in the art, so long as such modifications do not significantly inhibit the activity of the ribozyme.

Figure 7 shows an example of a Zinzyme A ribozyme motif that is chemically stabilized (see for example Beigelman *et al.*, International PCT publication No. WO 99/55857 and US Patent Application Serial No. 09/918,728).

10 Figure 8 shows an example of a DNAzyme motif described by Santoro *et al.*, 1997, *PNAS*, 94, 4262 and Joyce *et al.*, US 5,807,718 .

Figure 9 shows data demonstrating the inhibition of soluble VEGFR1 in a clinical study using ANGIOZYME (SEQ ID NO: 5977) .

15 Figure 10 shows an generalized outline for the mouse model of proliferative retinopathy showing the points of ribozyme administration.

Figure 11 shows a graph demonstrating the efficacy of a VEGF-receptor-targeted enzymatic nucleic acid molecule in a mouse model of proliferative retinopathy.

#### Detailed Description of the Invention

##### Nucleic Acid Molecules and Mechanism of Action

20 Enzymatic Nucleic Acid: Several varieties of naturally-occurring enzymatic nucleic acids are presently known. In addition, several *in vitro* selection (evolution) strategies (Orgel, 1979, *Proc. R. Soc. London*, B 205, 435) have been used to evolve new nucleic acid catalysts capable of catalyzing cleavage and ligation of phosphodiester linkages (Joyce, 1989, *Gene*, 82, 83-87; Beaudry *et al.*, 1992, *Science* 257, 635-641; Joyce, 1992, *Scientific American* 267, 25 90-97; Breaker *et al.*, 1994, *TIBTECH* 12, 268; Bartel *et al.*, 1993, *Science* 261:1411-1418; Szostak, 1993, *TIBS* 17, 89-93; Kumar *et al.*, 1995, *FASEB J.*, 9, 1183; Breaker, 1996, *Curr. Op. Biotech.*, 7, 442; Santoro *et al.*, 1997, *Proc. Natl. Acad. Sci.*, 94, 4262; Tang *et al.*, 1997, *RNA* 3, 914; Nakamaye & Eckstein, 1994, *supra*; Long & Uhlenbeck, 1994, *supra*; Ishizaka *et al.*, 1995, *supra*; Vaish *et al.*, 1997, *Biochemistry* 36, 6495; all of these are incorporated by 30 reference herein). Each can catalyze a series of reactions including the hydrolysis of

phosphodiester bonds in *trans* (and thus can cleave other nucleic acid molecules) under physiological conditions.

The enzymatic nature of an enzymatic nucleic acid molecule has significant advantages, one advantage being that the concentration of enzymatic nucleic acid molecule necessary to affect a therapeutic treatment is lower. This advantage reflects the ability of the enzymatic nucleic acid molecule to act enzymatically. Thus, a single enzymatic nucleic acid molecule is able to cleave many molecules of target nucleic acid. In addition, the enzymatic nucleic acid molecule is a highly specific inhibitor, with the specificity of inhibition depending not only on the base-pairing mechanism of binding to the target nucleic acid, but also on the mechanism of target nucleic acid cleavage. Single mismatches, or base-substitutions, near the site of cleavage can be chosen to completely eliminate catalytic activity of a enzymatic nucleic acid molecule.

Nucleic acid molecules having an endonuclease enzymatic activity are able to repeatedly cleave other separate nucleic acid molecules in a nucleotide base sequence-specific manner. With the proper design, such enzymatic nucleic acid molecules can be targeted to RNA transcripts, and achieve efficient cleavage *in vitro* (Zaug *et al.*, 324, *Nature* 429 1986; Uhlenbeck, 1987 *Nature* 328, 596; Kim *et al.*, 84 *Proc. Natl. Acad. Sci. USA* 8788, 1987; Dreyfus, 1988, *Einstein Quart. J. Bio. Med.*, 6, 92; Haseloff and Gerlach, 334 *Nature* 585, 1988; Cech, 260 *JAMA* 3030, 1988; and Jefferies *et al.*, 17 *Nucleic Acids Research* 1371, 1989; Santoro *et al.*, 1997 *supra*).

Because of their sequence specificity, *trans*-cleaving enzymatic nucleic acid molecules can be used as therapeutic agents for human disease (Usman & McSwiggen, 1995 *Ann. Rep. Med. Chem.* 30, 285-294; Christoffersen and Marr, 1995 *J. Med. Chem.* 38, 2023-2037). Enzymatic nucleic acid molecules can be designed to cleave specific nucleic acid targets within the background of cellular nucleic acid. Such a cleavage event renders the nucleic acid non-functional and abrogates protein expression from that nucleic acid. In this manner, synthesis of a protein associated with a disease state can be selectively inhibited (Warashina *et al.*, 1999, *Chemistry and Biology*, 6, 237-250).

Enzymatic nucleic acid molecules of the invention that are allosterically regulated ("allozymes") can be used to down-regulate VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2, expression. These allosteric enzymatic nucleic acids or allozymes (see for example Usman *et al.*, US Patent Application No. 09/877,526, George *et al.*, US Patent Nos. 5,834,186 and 5,741,679, Shih *et al.*, US Patent No. 5,589,332, Nathan *et al.*, US Patent No. 5,871,914, Nathan and Ellington, International PCT publication No. WO 00/24931, Breaker

*et al.*, International PCT Publication Nos. WO 00/26226 and 98/27104, and Sullenger *et al.*, US Patent Application Serial No. 09/205,520) are designed to respond to a signaling agent, for example, mutant VEGFR1 and/or VEGFR2 protein, wild-type VEGFR1 and/or VEGFR2 protein, mutant VEGFR1 and/or VEGFR2 RNA, wild-type VEGFR1 and/or VEGFR2 RNA, 5 other proteins and/or RNAs involved in VEGF signal transduction, compounds, metals, polymers, molecules and/or drugs that are targeted to VEGFR1 and/or VEGFR2 expression, which in turn modulates the activity of the enzymatic nucleic acid molecule. In response to interaction with a predetermined signaling agent, the activity of the allosteric enzymatic nucleic acid is activated or inhibited such that the expression of a particular target is 10 selectively down-regulated. The target can comprise wild-type VEGFR1 and/or VEGFR2, mutant VEGFR1 and/or VEGFR2, and/or a predetermined component of the VEGF signal transduction pathway. In a specific example, allosteric enzymatic nucleic acid molecules that are activated by interaction with a RNA encoding VEGF protein are used as therapeutic agents *in vivo*. The presence of RNA encoding the VEGF protein activates the allosteric 15 enzymatic nucleic acid molecule that subsequently cleaves the RNA encoding a VEGFR1 and/or VEGFR2 protein resulting in the inhibition of VEGFR1 and/or VEGFR2 protein expression.

In another non-limiting example, an allozyme can be activated by a VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 protein, peptide, or mutant polypeptide that causes 20 the allozyme to inhibit the expression of VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 genes, by, for example, cleaving RNA encoded by VEGF, VEGFR1 and/or VEGFR2 gene. In this non-limiting example, the allozyme acts as a decoy to inhibit the function of VEGF, VEGFR1 and/or VEGFR2 and also inhibit the expression of VEGF, VEGFR1 and/or VEGFR2 once activated by the VEGF, VEGFR1 and/or VEGFR2 protein.

25       Antisense: Antisense molecules can be modified or unmodified RNA, DNA, or mixed polymer oligonucleotides and primarily function by specifically binding to matching sequences resulting in inhibition of peptide synthesis (Wu-Pong, Nov 1994, *BioPharm*, 20-33). The antisense oligonucleotide binds to target RNA by Watson Crick base-pairing and blocks gene expression by preventing ribosomal translation of the bound sequences either by 30 steric blocking or by activating RNase H enzyme. Antisense molecules can also alter protein synthesis by interfering with RNA processing or transport from the nucleus into the cytoplasm (Mukhopadhyay & Roth, 1996, *Crit. Rev. in Oncogenesis* 7, 151-190).

In addition, binding of single stranded DNA to RNA can result in nuclease degradation of the heteroduplex (Wu-Pong, *supra*; Crooke, *supra*). To date, the only backbone modified

DNA chemistry which act as substrates for RNase H are phosphorothioates, phosphorodithioates, and borontrifluoridates. Recently it has been reported that 2'-arabino and 2'-fluoro arabino- containing oligos can also activate RNase H activity.

5 A number of antisense molecules have been described that utilize novel configurations of chemically modified nucleotides, secondary structure, and/or RNase H substrate domains (Woolf *et al.*, International PCT Publication No. WO 98/13526; Thompson *et al.*, International PCT Publication No. WO 99/54459; Hartmann *et al.*, USSN 60/101,174 which was filed on September 21, 1998) all of these are incorporated by reference herein in their entirety.

10 In addition, antisense deoxyoligoribonucleotides can be used to target RNA by means of DNA-RNA interactions, thereby activating RNase H, which digests the target RNA in the duplex. Antisense DNA can be expressed via the use of a single stranded DNA intracellular expression vector or equivalents and variations thereof.

15 Triplex Forming Oligonucleotides (TFO): Single stranded DNA can be designed to bind to genomic DNA in a sequence specific manner. TFOs are comprised of pyrimidine-rich oligonucleotides which bind DNA helices through Hoogsteen Base-pairing (Wu-Pong, *supra*). The resulting triple helix composed of the DNA sense, DNA antisense, and TFO disrupts RNA synthesis by RNA polymerase. The TFO mechanism can result in gene expression or cell death since binding can be irreversible (Mukhopadhyay & Roth, *supra*).

20 2-5A Antisense Chimera: The 2-5A system is an interferon mediated mechanism for RNA degradation found in higher vertebrates (Mitra *et al.*, 1996, *Proc Nat Acad Sci USA* 93, 6780-6785). Two types of enzymes, 2-5A synthetase and RNase L, are required for RNA cleavage. The 2-5A synthetases require double stranded RNA to form 2'-5' oligoadenylylates (2-5A). 2-5A then acts as an allosteric effector for utilizing RNase L which has the ability to 25 cleave single stranded RNA. The ability to form 2-5A structures with double stranded RNA makes this system particularly useful for inhibition of viral replication.

30 (2'-5') oligoadenylylate structures can be covalently linked to antisense molecules to form chimeric oligonucleotides capable of RNA cleavage (Torrence, *supra*). These molecules putatively bind and activate a 2-5A dependent RNase, the oligonucleotide/enzyme complex then binds to a target RNA molecule which can then be cleaved by the RNase enzyme.

RNAi: Double-stranded RNAs can suppress expression of homologous genes through an evolutionarily conserved process named RNA interference (RNAi) or post-transcriptional gene silencing (PTGS). One mechanism underlying silencing is the degradation of target mRNAs by an RNP complex, which contains short interfering RNAs (siRNAs) as guides to substrate selection. Short interfering RNAs are typically 21 to 23 nucleotides in length. A bidentate nuclease called Dicer has been implicated as the protein responsible for siRNA production. For example, a double-stranded RNA (dsRNA) matching a gene sequence is synthesized *in vitro* and introduced into a cell. The dsRNA feeds into a biological pathway and is broken into short pieces of short interfering (si) RNAs. With the help of cellular enzymes such as Dicer, the siRNA triggers the degradation of the messenger RNA that matches its sequence (see for example Tuschi *et al.*, International PCT Publication No. WO 01/75164; Bass, 2001, *Nature*, 411, 428-429; Elbashir *et al.*, 2001, *Nature*, 411, 494-498; and Kreutzer *et al.*, International PCT Publication No. WO 00/44895).

#### Target sites

Targets for useful nucleic acid molecules of the invention, such as enzymatic nucleic acid molecules, dsRNA, and antisense nucleic acids can be determined as disclosed in Draper *et al.*, WO 93/23569; Sullivan *et al.*, WO 93/23057; Thompson *et al.*, WO 94/02595; Draper *et al.*, WO 95/04818; McSwiggen *et al.*, US Patent No. 5,525,468, and hereby incorporated by reference herein in totality. Other examples include the following PCT applications, which concern inactivation of expression of disease-related genes: WO 95/23225, WO 95/13380, WO 94/02595, incorporated by reference herein. Rather than repeat the guidance provided in those documents here, below are provided specific examples of such methods, not limiting to those in the art. Enzymatic nucleic acid molecules and antisense to such targets are designed as described in those applications and synthesized to be tested *in vitro* and *in vivo*, as also described. The sequences of human VEGF, VEGFR1 and/or VEGFR2 RNAs are screened for optimal nucleic acid target sites using a computer-folding algorithm. Potential nucleic acid binding/cleavage sites are identified. While human sequences can be screened and nucleic acid molecules thereafter designed, as discussed in Stinchcomb *et al.*, WO 95/23225, mouse targeted enzymatic nucleic acid molecules can be useful to test efficacy of action of the nucleic acid molecule prior to testing in humans.

Nucleic acid molecule binding/cleavage sites are identified, for example enzymatic nucleic acid, antisense, and dsRNA mediated binding sites are chosen. For enzymatic nucleic acid molecules of the invention, the nucleic acid molecules are individually analyzed by computer folding (Jaeger *et al.*, 1989 *Proc. Natl. Acad. Sci. USA*, 86, 7706) to assess whether

the sequences fold into the appropriate secondary structure. Those nucleic acid molecules with unfavorable intramolecular interactions such as between the binding arms and the catalytic core can be eliminated from consideration. Varying binding arm lengths can be chosen to optimize activity.

5 Nucleic acids, such as antisense, RNAi, and/or enzymatic nucleic acid molecule binding/cleavage sites are identified and are designed to anneal to various sites in the nucleic acid target. The binding arms of enzymatic nucleic acid molecules of the invention are complementary to the target site sequences described above. Antisense and RNAi sequences are designed to have partial or complete complementarity to the nucleic acid target. The  
10 nucleic acid molecules can be chemically synthesized. The method of synthesis used follows the procedure for normal DNA/RNA synthesis as described below and in Usman *et al.*, 1987 *J. Am. Chem. Soc.*, 109, 7845; Scaringe *et al.*, 1990 *Nucleic Acids Res.*, 18, 5433; and Wincott *et al.*, 1995 *Nucleic Acids Res.* 23, 2677-2684; Caruthers *et al.*, 1992, *Methods in Enzymology* 211,3-19.

15 **Synthesis of Nucleic acid Molecules**

Synthesis of nucleic acids greater than 100 nucleotides in length is difficult using automated methods, and the therapeutic cost of such molecules is prohibitive. In this invention, small nucleic acid motifs ("small refers to nucleic acid motifs less than about 100 nucleotides in length, preferably less than about 80 nucleotides in length, and more preferably  
20 less than about 50 nucleotides in length; e.g., antisense oligonucleotides, enzymatic nucleic acids, aptamers, allozymes, decoys, siRNA etc.) are preferably used for exogenous delivery. The simple structure of these molecules increases the ability of the nucleic acid to invade targeted regions of RNA structure. Exemplary molecules of the instant invention are chemically synthesized, and others can similarly be synthesized.

25 DNA Oligonucleotides are synthesized using protocols known in the art as described in Caruthers *et al.*, 1992, *Methods in Enzymology* 211, 3-19, Thompson *et al.*, International PCT Publication No. WO 99/54459, Wincott *et al.*, 1995, *Nucleic Acids Res.* 23, 2677-2684, Wincott *et al.*, 1997, *Methods Mol. Bio.*, 74, 59, Brennan *et al.*, 1998, *Biotechnol Bioeng.*, 61, 33-45, and Brennan, US patent No. 6,001,311. All of these references are incorporated herein  
30 by reference. The synthesis of oligonucleotides makes use of common nucleic acid protecting and coupling groups, such as dimethoxytrityl at the 5'-end, and phosphoramidites at the 3'-end. In a non-limiting example, small scale syntheses are conducted on a 394 Applied Biosystems, Inc. synthesizer using a 0.2 μmol scale protocol with a 2.5 min coupling step for 2'-O-methylated nucleotides and a 45 sec coupling step for 2'-deoxy nucleotides. Table II

- outlines the amounts and the contact times of the reagents used in the synthesis cycle. Alternatively, syntheses at the 0.2  $\mu\text{mol}$  scale can be performed on a 96-well plate synthesizer, such as the instrument produced by Protogene (Palo Alto, CA) with minimal modification to the cycle. A 33-fold excess (60  $\mu\text{L}$  of 0.11 M = 6.6  $\mu\text{mol}$ ) of 2'-O-methyl phosphoramidite and a 105-fold excess of S-ethyl tetrazole (60  $\mu\text{L}$  of 0.25 M = 15  $\mu\text{mol}$ ) can be used in each coupling cycle of 2'-O-methyl residues relative to polymer-bound 5'-hydroxyl. A 22-fold excess (40  $\mu\text{L}$  of 0.11 M = 4.4  $\mu\text{mol}$ ) of deoxy phosphoramidite and a 70-fold excess of S-ethyl tetrazole (40  $\mu\text{L}$  of 0.25 M = 10  $\mu\text{mol}$ ) can be used in each coupling cycle of deoxy residues relative to polymer-bound 5'-hydroxyl. Average coupling yields on the 394 Applied Biosystems, Inc. synthesizer, determined by colorimetric quantitation of the trityl fractions, are typically 97.5-99%. Other oligonucleotide synthesis reagents for the 394 Applied Biosystems, Inc. synthesizer include; detritylation solution is 3% TCA in methylene chloride (ABI); capping is performed with 16% N-methyl imidazole in THF (ABI) and 10% acetic anhydride/10% 2,6-lutidine in THF (ABI); and oxidation solution is 16.9 mM  $\text{I}_2$ , 49 mM pyridine, 9% water in THF (PERSEPTIVE<sup>TM</sup>). Burdick & Jackson Synthesis Grade acetonitrile is used directly from the reagent bottle. S-Ethyltetrazole solution (0.25 M in acetonitrile) is made up from the solid obtained from American International Chemical, Inc. Alternately, for the introduction of phosphorothioate linkages, Beaucage reagent (3H-1,2-Benzodithiol-3-one 1,1-dioxide, 0.05 M in acetonitrile) is used.
- Deprotection of the DNA polynucleotides is performed as follows: the polymer-bound trityl-on oligoribonucleotide is transferred to a 4 mL glass screw top vial and suspended in a solution of 40% aq. methylamine (1 mL) at 65 °C for 10 min. After cooling to -20 °C, the supernatant is removed from the polymer support. The support is washed three times with 1.0 mL of EtOH:MeCN:H<sub>2</sub>O/3:1:1, vortexed and the supernatant is then added to the first supernatant. The combined supernatants, containing the oligoribonucleotide, are dried to a white powder.

The method of synthesis used for RNA oligonucleotides including certain nucleic acid molecules of the invention follows the procedure as described in Usman *et al.*, 1987, *J. Am. Chem. Soc.*, 109, 7845; Scaringe *et al.*, 1990, *Nucleic Acids Res.*, 18, 5433; and Wincott *et al.*, 1995, *Nucleic Acids Res.*, 23, 2677-2684 Wincott *et al.*, 1997, *Methods Mol. Bio.*, 74, 59, and makes use of common nucleic acid protecting and coupling groups, such as dimethoxytrityl at the 5'-end, and phosphoramidites at the 3'-end. In a non-limiting example, small scale syntheses are conducted on a 394 Applied Biosystems, Inc. synthesizer using a 0.2  $\mu\text{mol}$  scale protocol with a 7.5 min coupling step for alkylsilyl protected nucleotides and a 2.5 min coupling step for 2'-O-methylated nucleotides. Table II outlines the amounts and the

- contact times of the reagents used in the synthesis cycle. Alternatively, syntheses at the 0.2  $\mu\text{mol}$  scale can be done on a 96-well plate synthesizer, such as the instrument produced by Protogene (Palo Alto, CA) with minimal modification to the cycle. A 33-fold excess (60  $\mu\text{L}$  of 0.11 M = 6.6  $\mu\text{mol}$ ) of 2'-O-methyl phosphoramidite and a 75-fold excess of S-ethyl tetrazole (60  $\mu\text{L}$  of 0.25 M = 15  $\mu\text{mol}$ ) can be used in each coupling cycle of 2'-O-methyl residues relative to polymer-bound 5'-hydroxyl. A 66-fold excess (120  $\mu\text{L}$  of 0.11 M = 13.2  $\mu\text{mol}$ ) of alkylsilyl (ribo) protected phosphoramidite and a 150-fold excess of S-ethyl tetrazole (120  $\mu\text{L}$  of 0.25 M = 30  $\mu\text{mol}$ ) can be used in each coupling cycle of ribo residues relative to polymer-bound 5'-hydroxyl. Average coupling yields on the 394 Applied Biosystems, Inc. synthesizer, determined by colorimetric quantitation of the trityl fractions, are typically 97.5-99%. Other oligonucleotide synthesis reagents for the 394 Applied Biosystems, Inc. synthesizer include; detritylation solution is 3% TCA in methylene chloride (ABI); capping is performed with 16% N-methyl imidazole in THF (ABI) and 10% acetic anhydride/10% 2,6-lutidine in THF (ABI); oxidation solution is 16.9 mM I<sub>2</sub>, 49 mM pyridine, 9% water in THF (PERSEPTIVE™). Burdick & Jackson Synthesis Grade acetonitrile is used directly from the reagent bottle. S-Ethyltetrazole solution (0.25 M in acetonitrile) is made up from the solid obtained from American International Chemical, Inc. Alternately, for the introduction of phosphorothioate linkages, Beaucage reagent (3H-1,2-Benzodithiol-3-one 1,1-dioxide 0.05 M in acetonitrile) is used.
- Deprotection of the RNA is performed using either a two-pot or one-pot protocol. For the two-pot protocol, the polymer-bound trityl-on oligoribonucleotide is transferred to a 4 mL glass screw top vial and suspended in a solution of 40% aq. methylamine (1 mL) at 65 °C for 10 min. After cooling to -20 °C, the supernatant is removed from the polymer support. The support is washed three times with 1.0 mL of EtOH:MeCN:H<sub>2</sub>O/3:1:1, vortexed and the supernatant is then added to the first supernatant. The combined supernatants, containing the oligoribonucleotide, are dried to a white powder. The base deprotected oligoribonucleotide is resuspended in anhydrous TEA/HF/NMP solution (300  $\mu\text{L}$  of a solution of 1.5 mL N-methylpyrrolidinone, 750  $\mu\text{L}$  TEA and 1 mL TEA•3HF to provide a 1.4 M HF concentration) and heated to 65 °C. After 1.5 h, the oligomer is quenched with 1.5 M NH<sub>4</sub>HCO<sub>3</sub>.
- Alternatively, for the one-pot protocol, the polymer-bound trityl-on oligoribonucleotide is transferred to a 4 mL glass screw top vial and suspended in a solution of 33% ethanolic methylamine/DMSO: 1/1 (0.8 mL) at 65 °C for 15 min. The vial is brought to r.t. TEA•3HF (0.1 mL) is added and the vial is heated at 65 °C for 15 min. The sample is cooled at -20 °C and then quenched with 1.5 M NH<sub>4</sub>HCO<sub>3</sub>.

For purification of the trityl-on oligomers, the quenched NH<sub>4</sub>HCO<sub>3</sub> solution is loaded onto a C-18 containing cartridge that had been prewashed with acetonitrile followed by 50 mM TEAA. After washing the loaded cartridge with water, the RNA is detritylated with 0.5% TFA for 13 min. The cartridge is then washed again with water, salt exchanged with 1 M NaCl and washed with water again. The oligonucleotide is then eluted with 30% acetonitrile.

Inactive hammerhead ribozymes or binding attenuated control (BAC) oligonucleotides are synthesized by substituting a U for G<sub>5</sub> and a U for A<sub>14</sub> (numbering from Hertel, K. J., *et al.*, 1992, *Nucleic Acids Res.*, 20, 3252). Similarly, one or more nucleotide substitutions can be introduced in other enzymatic nucleic acid molecules to inactivate the molecule and such molecules can serve as a negative control.

The average stepwise coupling yields are typically >98% (Wincott *et al.*, 1995 *Nucleic Acids Res.* 23, 2677-2684). Those of ordinary skill in the art will recognize that the scale of synthesis can be adapted to be larger or smaller than the example described above including but not limited to 96 well format, all that is important is the ratio of chemicals used in the reaction.

Alternatively, the nucleic acid molecules of the present invention can be synthesized separately and joined together post-synthetically, for example by ligation (Moore *et al.*, 1992, *Science* 256, 9923; Draper *et al.*, International PCT publication No. WO 93/23569; Shabarova *et al.*, 1991, *Nucleic Acids Research* 19, 4247; Bellon *et al.*, 1997, *Nucleosides & Nucleotides*, 16, 951; Bellon *et al.*, 1997, *Bioconjugate Chem.* 8, 204).

Preferably, the nucleic acid molecules of the present invention are modified extensively to enhance stability by modification with nuclease resistant groups, for example, 2'-amino, 2'-C-allyl, 2'-flouro, 2'-O-methyl, 2'-H (for a review see Usman and Cedergren, 1992, *TIBS* 17, 34; Usman *et al.*, 1994, *Nucleic Acids Symp. Ser.* 31, 163). Ribozymes are purified by gel electrophoresis using general methods or are purified by high pressure liquid chromatography (HPLC; See Wincott *et al.*, Supra, the totality of which is hereby incorporated herein by reference) and are re-suspended in water.

Optimizing Activity of the nucleic acid molecule of the invention.

Chemically synthesizing nucleic acid molecules with modifications (base, sugar and/or phosphate) that prevent their degradation by serum ribonucleases can increase their potency (see e.g., Eckstein *et al.*, International Publication No. WO 92/07065; Perrault *et al.*, 1990 *Nature* 344, 565; Pieken *et al.*, 1991, *Science* 253, 314; Usman and Cedergren, 1992, *Trends*

in *Biochem. Sci.* 17, 334; Usman *et al.*, International Publication No. WO 93/15187; and Rossi *et al.*, International Publication No. WO 91/03162; Sproat, US Patent No. 5,334,711; Gold *et al.*, US 6,300,074; and Burgin *et al.*, *supra*; all of which are incorporated by reference herein). Modifications which enhance their efficacy in cells, and removal of bases from 5 nucleic acid molecules to shorten oligonucleotide synthesis times and reduce chemical requirements are desired. (All these publications are hereby incorporated by reference herein).

There are several examples in the art describing sugar, base and phosphate modifications that can be introduced into nucleic acid molecules with significant 10 enhancement in their nuclease stability and efficacy. For example, oligonucleotides are modified to enhance stability and/or enhance biological activity by modification with nuclease resistant groups, for example, 2'-amino, 2'-C-allyl, 2'-flouro, 2'-O-methyl, 2'-H, nucleotide base modifications (for a review see Usman and Cedergren, 1992, *TIBS*, 17, 34; Usman *et al.*, 1994, *Nucleic Acids Symp. Ser.* 31, 163; Burgin *et al.*, 1996, *Biochemistry*, 35, 15 14090). Sugar modification of nucleic acid molecules have been extensively described in the art (see Eckstein *et al.*, *International Publication* PCT No. WO 92/07065; Perrault *et al.* *Nature*, 1990, 344, 565-568; Pieken *et al.* *Science*, 1991, 253, 314-317; Usman and Cedergren, *Trends in Biochem. Sci.*, 1992, 17, 334-339; Usman *et al.* *International Publication* PCT No. WO 93/15187; Sproat, US Patent No. 5,334,711 and Beigelman *et al.*, 20 1995, *J. Biol. Chem.*, 270, 25702; Beigelman *et al.*, *International PCT publication* No. WO 97/26270; Beigelman *et al.*, US Patent No. 5,716,824; Usman *et al.*, US patent No. 5,627,053; Woolf *et al.*, *International PCT Publication* No. WO 98/13526; Thompson *et al.*, USSN 60/082,404 which was filed on April 20, 1998; Karpeisky *et al.*, 1998, *Tetrahedron Lett.*, 39, 1131; Earnshaw and Gait, 1998, *Biopolymers (Nucleic acid Sciences)*, 48, 39-55; 25 Verma and Eckstein, 1998, *Annu. Rev. Biochem.*, 67, 99-134; and Burlina *et al.*, 1997, *Bioorg. Med. Chem.*, 5, 1999-2010; all of the references are hereby incorporated in their totality by reference herein). Such publications describe general methods and strategies to determine the location of incorporation of sugar, base and/or phosphate modifications and the like into ribozymes without inhibiting catalysis, and are incorporated by reference herein. In 30 view of such teachings, similar modifications can be used as described herein to modify the nucleic acid molecules of the instant invention.

While chemical modification of oligonucleotide internucleotide linkages with phosphorothioate, phosphorothioate, and/or 5'-methylphosphonate linkages improves 35 stability, too many of these modifications can cause some toxicity. Therefore when designing nucleic acid molecules the amount of these internucleotide linkages should be minimized.

The reduction in the concentration of these linkages should lower toxicity resulting in increased efficacy and higher specificity of these molecules.

- Nucleic acid molecules having chemical modifications that maintain or enhance activity are provided. Such nucleic acid is also generally more resistant to nucleases than unmodified 5 nucleic acid. Thus, in a cell and/or *in vivo* the activity may not be significantly lowered. Therapeutic nucleic acid molecules delivered exogenously are optimally stable within cells until translation of the target RNA has been inhibited long enough to reduce the levels of the undesirable protein. This period of time varies between hours to days depending upon the disease state. Clearly, nucleic acid molecules must be resistant to nucleases in order to 10 function as effective intracellular therapeutic agents. Improvements in the chemical synthesis of RNA and DNA (Wincott *et al.*, 1995 *Nucleic Acids Res.* 23, 2677; Caruthers *et al.*, 1992, *Methods in Enzymology* 211, 3-19 (incorporated by reference herein) have expanded the ability to modify nucleic acid molecules by introducing nucleotide modifications to enhance their nuclease stability as described above.
- 15 In one embodiment, nucleic acid molecules of the invention include one or more G-clamp nucleotides. A G-clamp nucleotide is a modified cytosine analog wherein the modifications confer the ability to hydrogen bond both Watson-Crick and Hoogsteen faces of a complementary guanine within a duplex, see for example Lin and Matteucci, 1998, *J. Am. Chem. Soc.*, 120, 8531-8532. A single G-clamp analog substitution within an oligonucleotide 20 can result in substantially enhanced helical thermal stability and mismatch discrimination when hybridized to complementary oligonucleotides. The inclusion of such nucleotides in nucleic acid molecules of the invention results in both enhanced affinity and specificity to nucleic acid targets. In another embodiment, nucleic acid molecules of the invention include 25 one or more LNA "locked nucleic acid" nucleotides such as a 2', 4'-C methylene bicyclo nucleotide (see for example Wengel *et al.*, International PCT Publication No. WO 00/66604 and WO 99/14226).

- In another embodiment, the invention features conjugates and/or complexes of nucleic acid molecules targeting VEGF receptors such as VEGFR1 and/or VEGFR2. Such conjugates and/or complexes can be used to facilitate delivery of molecules into a biological 30 system, such as cells. The conjugates and complexes provided by the instant invention can impart therapeutic activity by transferring therapeutic compounds across cellular membranes, altering the pharmacokinetics, and/or modulating the localization of nucleic acid molecules of the invention. The present invention encompasses the design and synthesis of novel conjugates and complexes for the delivery of molecules, including but not limited to small

molecules, lipids, phospholipids, nucleosides, nucleotides, nucleic acids, antibodies, toxins, negatively charged polymers and other polymers, for example proteins, peptides, hormones, carbohydrates, polyethylene glycols, or polyamines, across cellular membranes. In general, the transporters described are designed to be used either individually or as part of a multi-  
5 component system, with or without degradable linkers. These compounds are expected to improve delivery and/or localization of nucleic acid molecules of the invention into a number of cell types originating from different tissues, in the presence or absence of serum (see Sullenger and Cech, US 5,854,038). Conjugates of the molecules described herein can be attached to biologically active molecules via linkers that are biodegradable, such as  
10 biodegradable nucleic acid linker molecules.

The term "biodegradable nucleic acid linker molecule" as used herein, refers to a nucleic acid molecule that is designed as a biodegradable linker to connect one molecule to another molecule, for example, a biologically active molecule. The stability of the biodegradable nucleic acid linker molecule can be modulated by using various combinations  
15 of ribonucleotides, deoxyribonucleotides, and chemically modified nucleotides, for example, 2'-O-methyl, 2'-fluoro, 2'-amino, 2'-O-amino, 2'-C-allyl, 2'-O-allyl, and other 2'-modified or base modified nucleotides. The biodegradable nucleic acid linker molecule can be a dimer, trimer, tetramer or longer nucleic acid molecule, for example, an oligonucleotide of about 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides in length, or can  
20 comprise a single nucleotide with a phosphorus based linkage, for example, a phosphoramidate or phosphodiester linkage. The biodegradable nucleic acid linker molecule can also comprise nucleic acid backbone, nucleic acid sugar, or nucleic acid base modifications.

The term "biodegradable" as used herein, refers to degradation in a biological system,  
25 for example enzymatic degradation or chemical degradation.

The term "biologically active molecule" as used herein, refers to compounds or molecules that are capable of eliciting or modifying a biological response in a system. Non-limiting examples of biologically active molecules contemplated by the instant invention include therapeutically active molecules such as antibodies, hormones, antivirals, peptides, proteins, therapeutics, small molecules, vitamins, co-factors, nucleosides, nucleotides, oligonucleotides, enzymatic nucleic acids, antisense nucleic acids, triplex forming oligonucleotides, 2,5-A chimeras, siRNA, dsRNA, allozymes, aptamers, decoys and analogs thereof. Biologically active molecules of the invention also include molecules capable of modulating the pharmacokinetics and/or pharmacodynamics of other biologically active

molecules, for example, lipids and polymers such as polyamines, polyamides, polyethylene glycol and other polyethers.

The term "phospholipid" as used herein, refers to a hydrophobic molecule comprising at least one phosphorus group. For example, a phospholipid can comprise a phosphorus containing group and saturated or unsaturated alkyl group, optionally substituted with OH, COOH, oxo, amine, or substituted or unsubstituted aryl groups.

Therapeutic nucleic acid molecules (e.g., enzymatic nucleic acid molecules and antisense nucleic acid molecules) delivered exogenously are optimally stable within cells until translation of the target RNA has been inhibited long enough to reduce the levels of the undesirable protein. This period of time varies between hours to days depending upon the disease state. These nucleic acid molecules should be resistant to nucleases in order to function as effective intracellular therapeutic agents. Improvements in the chemical synthesis of nucleic acid molecules described in the instant invention and in the art have expanded the ability to modify nucleic acid molecules by introducing nucleotide modifications to enhance their nuclease stability as described above.

In another embodiment, nucleic acid catalysts having chemical modifications that maintain or enhance enzymatic activity are provided. Such nucleic acids are also generally more resistant to nucleases than unmodified nucleic acid. Thus, in a cell and/or *in vivo* the activity of the nucleic acid may not be significantly lowered. As exemplified herein such enzymatic nucleic acids are useful in a cell and/or *in vivo* even if activity over all is reduced 10 fold (Burgin *et al.*, 1996, *Biochemistry*, 35, 14090). Such enzymatic nucleic acids herein are said to "maintain" the enzymatic activity of an all RNA ribozyme or all DNA DNazyme.

In another aspect the nucleic acid molecules comprise a 5' and/or a 3'- cap structure.

By "cap structure" is meant chemical modifications, which have been incorporated at either terminus of the oligonucleotide (see for example Wincott *et al.*, WO 97/26270, incorporated by reference herein). These terminal modifications protect the nucleic acid molecule from exonuclease degradation, and can help in delivery and/or localization within a cell. The cap can be present at the 5'-terminus (5'-cap) or at the 3'-terminus (3'-cap) or can be present on both terminus. In non-limiting examples, the 5'-cap includes inverted abasic residue (moiety), 4',5'-methylene nucleotide; 1-(beta-D-erythrofuranosyl) nucleotide, 4'-thio nucleotide, carbocyclic nucleotide; 1,5-anhydrohexitol nucleotide; L-nucleotides; alpha-nucleotides; modified base nucleotide; phosphorodithioate linkage; *threo*-pentofuranosyl nucleotide; acyclic 3',4'-seco nucleotide; acyclic 3,4-dihydroxybutyl nucleotide; acyclic 3,5-

dihydroxypentyl nucleotide, 3'-3'-inverted nucleotide moiety; 3'-3'-inverted abasic moiety; 3'-2'-inverted nucleotide moiety; 3'-2'-inverted abasic moiety; 1,4-butanediol phosphate; 3'-phosphoramidate; hexylphosphate; aminohexyl phosphate; 3'-phosphate; 3'-phosphorothioate; phosphorodithioate; or bridging or non-bridging methylphosphonate moiety (for more details see Wincott *et al.*, International PCT publication No. WO 97/26270, incorporated by reference herein).

In another embodiment the 3'-cap includes, for example 4',5'-methylene nucleotide; 1-(beta-D-erythrofuranosyl) nucleotide; 4'-thio nucleotide, carbocyclic nucleotide; 5'-amino-alkyl phosphate; 1,3-diamino-2-propyl phosphate, 3-aminopropyl phosphate; 6-aminohexyl phosphate; 1,2-aminododecyl phosphate; hydroxypropyl phosphate; 1,5-anhydrohexitol nucleotide; L-nucleotide; alpha-nucleotide; modified base nucleotide; phosphorodithioate; *threo*-pentofuranosyl nucleotide; acyclic 3',4'-seco nucleotide; 3,4-dihydroxybutyl nucleotide; 3,5-dihydroxypentyl nucleotide, 5'-5'-inverted nucleotide moiety; 5'-5'-inverted abasic moiety; 5'-phosphoramidate; 5'-phosphorothioate; 1,4-butanediol phosphate; 5'-amino; bridging and/or non-bridging 5'-phosphoramidate, phosphorothioate and/or phosphorodithioate, bridging or non bridging methylphosphonate and 5'-mercapto moieties (for more details see Beaucage and Iyer, 1993, *Tetrahedron* 49, 1925; incorporated by reference herein).

By the term "non-nucleotide" is meant any group or compound which can be incorporated into a nucleic acid chain in the place of one or more nucleotide units, including either sugar and/or phosphate substitutions, and allows the remaining bases to exhibit their enzymatic activity. The group or compound is abasic in that it does not contain a commonly recognized nucleotide base, such as adenosine, guanine, cytosine, uracil or thymine.

An "alkyl" group refers to a saturated aliphatic hydrocarbon, including straight-chain, branched-chain, and cyclic alkyl groups. Preferably, the alkyl group has 1 to 12 carbons. More preferably it is a lower alkyl of from 1 to 7 carbons, more preferably 1 to 4 carbons. The alkyl group can be substituted or unsubstituted. When substituted the substituted group(s) is preferably, hydroxyl, cyano, alkoxy, =O, =S, NO<sub>2</sub> or N(CH<sub>3</sub>)<sub>2</sub>, amino, or SH. The term also includes alkenyl groups which are unsaturated hydrocarbon groups containing at least one carbon-carbon double bond, including straight-chain, branched-chain, and cyclic groups. Preferably, the alkenyl group has 1 to 12 carbons. More preferably it is a lower alkenyl of from 1 to 7 carbons, more preferably 1 to 4 carbons. The alkenyl group can be substituted or unsubstituted. When substituted the substituted group(s) is preferably, hydroxyl, cyano, alkoxy, =O, =S, NO<sub>2</sub>, halogen, N(CH<sub>3</sub>)<sub>2</sub>, amino, or SH. The term "alkyl" also includes alkynyl groups which have an unsaturated hydrocarbon group containing at least

one carbon-carbon triple bond, including straight-chain, branched-chain, and cyclic groups. Preferably, the alkynyl group has 1 to 12 carbons. More preferably it is a lower alkynyl of from 1 to 7 carbons, more preferably 1 to 4 carbons. The alkynyl group can be substituted or unsubstituted. When substituted the substituted group(s) is preferably, hydroxyl, cyano, 5 alkoxy, =O, =S, NO<sub>2</sub> or N(CH<sub>3</sub>)<sub>2</sub>, amino or SH.

Such alkyl groups can also include aryl, alkylaryl, carbocyclic aryl, heterocyclic aryl, amide and ester groups. An "aryl" group refers to an aromatic group which has at least one ring having a conjugated p electron system and includes carbocyclic aryl, heterocyclic aryl and biaryl groups, all of which can be optionally substituted. The preferred substituent(s) of 10 aryl groups are halogen, trihalomethyl, hydroxyl, SH, OH, cyano, alkoxy, alkyl, alkenyl, alkynyl, and amino groups. An "alkylaryl" group refers to an alkyl group (as described above) covalently joined to an aryl group (as described above). Carbocyclic aryl groups are groups wherein the ring atoms on the aromatic ring are all carbon atoms. The carbon atoms are optionally substituted. Heterocyclic aryl groups are groups having from 1 to 3 15 heteroatoms as ring atoms in the aromatic ring and the remainder of the ring atoms are carbon atoms. Suitable heteroatoms include oxygen, sulfur, and nitrogen, and include furanyl, thienyl, pyridyl, pyrrolyl, N-lower alkyl pyrrolo, pyrimidyl, pyrazinyl, imidazolyl and the like, all optionally substituted. An "amide" refers to an -C(O)-NH-R, where R is either alkyl, aryl, alkylaryl or hydrogen. An "ester" refers to an -C(O)-OR', where R is either alkyl, aryl, 20 alkylaryl or hydrogen.

By "nucleotide" is meant a heterocyclic nitrogenous base in N-glycosidic linkage with a phosphorylated sugar. Nucleotides are recognized in the art to include natural bases (standard), and modified bases well known in the art. Such bases are generally located at the 1' position of a nucleotide sugar moiety. Nucleotides generally comprise a base, sugar and a 25 phosphate group. The nucleotides can be unmodified or modified at the sugar, phosphate and/or base moiety, (also referred to interchangeably as nucleotide analogs, modified nucleotides, non-natural nucleotides, non-standard nucleotides and other; see for example, Usman and McSwiggen, *supra*; Eckstein *et al.*, International PCT Publication No. WO 92/07065; Usman *et al.*, International PCT Publication No. WO 93/15187; Uhlman & 30 Peyman, *supra* all are hereby incorporated by reference herein). There are several examples of modified nucleic acid bases known in the art as summarized by Limbach *et al.*, 1994, Nucleic Acids Res. 22, 2183. Some of the non-limiting examples of chemically modified and other natural nucleic acid bases that can be introduced into nucleic acids include, for example, inosine, purine, pyridin-4-one, pyridin-2-one, phenyl, pseudouracil, 2, 4, 6-trimethoxy 35 benzene, 3-methyl uracil, dihydrouridine, naphthyl, aminophenyl, 5-alkylcytidines (e.g.,

- 5-methylcytidine), 5-alkyluridines (e.g., ribothymidine), 5-halouridine (e.g., 5-bromouridine) or 6-azapyrimidines or 6-alkylpyrimidines (e.g. 6-methyluridine), propyne, quenosine, 2-thiouridine, 4-thiouridine, wybutoxine, wybutoxosine, 4-acetylcytidine, 5-(carboxyhydroxymethyl)uridine, 5'-carboxymethylaminomethyl-2-thiouridine, 5-
- 5 carboxymethylaminomethyluridine, beta-D-galactosylqueosine, 1-methyladenosine, 1-methylinosine, 2,2-dimethylguanosine, 3-methylcytidine, 2-methyladenosine, 2-methylguanosine, N6-methyladenosine, 7-methylguanosine, 5-methoxyaminomethyl-2-thiouridine, 5-methylaminomethyluridine, 5-methylcarboxymethyluridine, 5-methoxyuridine, 5-methyl-2-thiouridine, 2-methylthio-N6-isopentenyladenosine, beta-D-mannosylqueosine, uridine-5-oxyacetic acid, 2-thiocytidine, threonine derivatives and others (Burgin *et al.*, 1996, Biochemistry, 35, 14090; Uhlman & Peyman, *supra*). By "modified bases" in this aspect is meant nucleotide bases other than adenine, guanine, cytosine and uracil at 1' position or their equivalents; such bases can be used at any position, for example, within the catalytic core of an enzymatic nucleic acid molecule and/or in the substrate-binding regions of the nucleic acid molecule.

By "nucleoside" is meant a heterocyclic nitrogenous base in N-glycosidic linkage with a sugar. Nucleosides are recognized in the art to include natural bases (standard), and modified bases well known in the art. Such bases are generally located at the 1' position of a nucleoside sugar moiety. Nucleosides generally comprise a base and sugar group. The nucleosides can be unmodified or modified at the sugar, and/or base moiety, (also referred to interchangeably as nucleoside analogs, modified nucleosides, non-natural nucleosides, non-standard nucleosides and other; see for example, Usman and McSwiggen, *supra*; Eckstein *et al.*, International PCT Publication No. WO 92/07065; Usman *et al.*, International PCT Publication No. WO 93/15187; Uhlman & Peyman, *supra* all are hereby incorporated by reference herein). There are several examples of modified nucleic acid bases known in the art as summarized by Limbach *et al.*, 1994, Nucleic Acids Res. 22, 2183. Some of the non-limiting examples of chemically modified and other natural nucleic acid bases that can be introduced into nucleic acids include, inosine, purine, pyridin-4-one, pyridin-2-one, phenyl, pseudouracil, 2, 4, 6-trimethoxy benzene, 3-methyl uracil, dihydrouridine, naphthyl, aminophenyl, 5-alkylcytidines (e.g., 5-methylcytidine), 5-alkyluridines (e.g., ribothymidine), 5-halouridine (e.g., 5-bromouridine) or 6-azapyrimidines or 6-alkylpyrimidines (e.g. 6-methyluridine), propyne, quenosine, 2-thiouridine, 4-thiouridine, wybutoxine, wybutoxosine, 4-acetylcytidine, 5-(carboxyhydroxymethyl)uridine, 5'-carboxymethylaminomethyl-2-thiouridine, 5-carboxymethylaminomethyluridine, beta-D-galactosylqueosine, 1-methyladenosine, 1-methylinosine, 2,2-dimethylguanosine, 3-methylcytidine, 2-methyladenosine, 2-methylguanosine, N6-methyladenosine, 7-methylguanosine, 5-

methoxyaminomethyl-2-thiouridine, 5-methylaminomethyluridine, 5-methylcarbonylmethyluridine, 5-methoxyuridine, 5-methyl-2-thiouridine, 2-methylthio-N6-isopentenyladenosine, beta-D-mannosylqueosine, uridine-5-oxyacetic acid, 2-thiocytidine, threonine derivatives and others (Burgin *et al.*, 1996, *Biochemistry*, 35, 14090; Uhlman & 5 Peyman, *supra*). By "modified bases" in this aspect is meant nucleoside bases other than adenine, guanine, cytosine and uracil at 1' position or their equivalents; such bases can be used at any position, for example, within the catalytic core of an enzymatic nucleic acid molecule and/or in the substrate-binding regions of the nucleic acid molecule.

In one embodiment, the invention features modified enzymatic nucleic acid molecules 10 with phosphate backbone modifications comprising one or more phosphorothioate, phosphorodithioate, methylphosphonate, morpholino, amidate carbamate, carboxymethyl, acetamidate, polyamide, sulfonate, sulfonamide, sulfamate, formacetal, thioformacetal, and/or alkylsilyl, substitutions. For a review of oligonucleotide backbone modifications see Hunziker and Leumann, 1995, *Nucleic Acid Analogues: Synthesis and Properties*, in *Modern 15 Synthetic Methods*, VCH, 331-417, and Mesmaeker *et al.*, 1994, *Novel Backbone Replacements for Oligonucleotides*, in *Carbohydrate Modifications in Antisense Research*, ACS, 24-39. These references are hereby incorporated by reference herein.

By "abasic" is meant sugar moieties lacking a base or having other chemical groups in place of a base at the 1' position, for example a 3',3'-linked or 5',5'-linked deoxyabasic 20 ribose derivative (for more details see Wincott *et al.*, International PCT publication No. WO 97/26270).

By "unmodified nucleoside" is meant one of the bases adenine, cytosine, guanine, thymine, uracil joined to the 1' carbon of β-D-ribo-furanose.

By "modified nucleoside" is meant any nucleotide base which contains a modification 25 in the chemical structure of an unmodified nucleotide base, sugar and/or phosphate.

In connection with 2'-modified nucleotides as described for the present invention, by "amino" is meant 2'-NH<sub>2</sub> or 2'-O- NH<sub>2</sub>, which can be modified or unmodified. Such modified groups are described, for example, in Eckstein *et al.*, U.S. Patent 5,672,695 and Matulic-Adamic *et al.*, WO 98/28317, respectively, which are both incorporated by reference 30 in their entireties.

Various modifications to nucleic acid (e.g., antisense and ribozyme) structure can be made to enhance the utility of these molecules. For example, such modifications can enhance

shelf-life, half-life *in vitro*, stability, and ease of introduction of such oligonucleotides to the target site, including, *e.g.*, enhancing penetration of cellular membranes and conferring the ability to recognize and bind to targeted cells.

Use of the nucleic acid-based molecules of the invention can lead to better treatment of the disease progression by affording the possibility of combination therapies (*e.g.*, multiple enzymatic nucleic acid molecules targeted to different genes, enzymatic nucleic acid molecules coupled with known small molecule inhibitors, or intermittent treatment with combinations of enzymatic nucleic acid molecules (including different enzymatic nucleic acid molecule motifs) and/or other chemical or biological molecules). The treatment of patients with nucleic acid molecules can also include combinations of different types of nucleic acid molecules. Therapies can be devised which include a mixture of enzymatic nucleic acid molecules (including different enzymatic nucleic acid molecule motifs), allozymes, antisense, dsRNA, aptamers, and/or 2-5A chimera molecules to one or more targets to alleviate symptoms of a disease.

15 **Administration of Nucleic Acid Molecules**

Methods for the delivery of nucleic acid molecules are described in Akhtar *et al.*, 1992, *Trends Cell Bio.*, 2, 139; and *Delivery Strategies for Antisense Oligonucleotide Therapeutics*, ed. Akhtar, 1995 which are both incorporated herein by reference. Sullivan *et al.*, PCT WO 94/02595, further describes the general methods for delivery of enzymatic RNA molecules. These protocols can be utilized for the delivery of virtually any nucleic acid molecule. Nucleic acid molecules can be administered to cells by a variety of methods known to those familiar to the art, including, but not restricted to, encapsulation in liposomes, by iontophoresis, or by incorporation into other vehicles, such as hydrogels, cyclodextrins, biodegradable nanocapsules, and bioadhesive microspheres. Alternatively, the nucleic acid/vehicle combination is locally delivered by direct injection or by use of an infusion pump. Other routes of delivery include, but are not limited to oral (tablet or pill form) and/or intrathecal delivery (Gold, 1997, *Neuroscience*, 76, 1153-1158). Other approaches include the use of various transport and carrier systems, for example though the use of conjugates and biodegradable polymers. For a comprehensive review on drug delivery strategies including CNS delivery, see Ho *et al.*, 1999, *Curr. Opin. Mol. Ther.*, 1, 336-343 and Jain, *Drug Delivery Systems: Technologies and Commercial Opportunities*, Decision Resources, 1998 and Grootenhuis *et al.*, 1997, *J. NeuroVirol.*, 3, 387-400. More detailed descriptions of nucleic acid delivery and administration are provided in Sullivan *et al.*, *supra*, Draper *et al.*, PCT

WO93/23569, Beigelman *et al.*, PCT WO99/05094, and Klimuk *et al.*, PCT WO99/04819 all of which have been incorporated by reference herein.

The molecules of the instant invention can be used as pharmaceutical agents. Pharmaceutical agents prevent, inhibit the occurrence, or treat (alleviate a symptom to some extent, preferably all of the symptoms) of a disease state in a patient.

The polynucleotides of the invention can be administered (e.g., RNA, DNA or protein) and introduced into a patient by any standard means, with or without stabilizers, buffers, and the like, to form a pharmaceutical composition. When it is desired to use a liposome delivery mechanism, standard protocols for formation of liposomes can be followed. The compositions of the present invention can also be formulated and used as tablets, capsules or elixirs for oral administration; suppositories for rectal administration; sterile solutions; suspensions for injectable administration; and the other compositions known in the art.

The present invention also includes pharmaceutically acceptable formulations of the compounds described. These formulations include salts of the above compounds, e.g., acid addition salts, for example, salts of hydrochloric, hydrobromic, acetic acid, and benzene sulfonic acid.

A pharmacological composition or formulation refers to a composition or formulation in a form suitable for administration, e.g., systemic administration, into a cell or patient, preferably a human. Suitable forms, in part, depend upon the use or the route of entry, for example oral, transdermal, or by injection. Such forms should not prevent the composition or formulation from reaching a target cell (*i.e.*, a cell to which the negatively charged polymer is desired to be delivered to). For example, pharmacological compositions injected into the blood stream should be soluble. Other factors are known in the art, and include considerations such as toxicity and forms which prevent the composition or formulation from exerting its effect.

By "systemic administration" is meant *in vivo* systemic absorption or accumulation of drugs in the blood stream followed by distribution throughout the entire body. Administration routes which lead to systemic absorption include, without limitations: intravenous, subcutaneous, intraperitoneal, inhalation, oral, intrapulmonary and intramuscular. Each of these administration routes expose the desired negatively charged polymers, e.g., nucleic acids, to an accessible diseased tissue. The rate of entry of a drug into the circulation has been shown to be a function of molecular weight or size. The use of a liposome or other drug carrier comprising the compounds of the instant invention can

potentially localize the drug, for example, in certain tissue types, such as the tissues of the reticular endothelial system (RES). A liposome formulation which can facilitate the association of drug with the surface of cells, such as, lymphocytes and macrophages is also useful. This approach can provide enhanced delivery of the drug to target cells by taking  
5 advantage of the specificity of macrophage and lymphocyte immune recognition of abnormal cells, such as cells implicated in endometriosis, birth control, endometrial tumors, gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), menopausal dysfunction, and endometrial carcinoma.

By pharmaceutically acceptable formulation is meant, a composition or formulation that  
10 allows for the effective distribution of the nucleic acid molecules of the instant invention in the physical location most suitable for their desired activity. Non-limiting examples of agents suitable for formulation with the nucleic acid molecules of the instant invention include: PEG conjugated nucleic acids, phospholipid conjugated nucleic acids, nucleic acids containing lipophilic moieties, phosphorothioates, P-glycoprotein inhibitors (such as Pluronic P85) which can enhance entry of drugs into various tissues, for example the CNS (Jollet-Riant and Tillement, 1999, *Fundam. Clin. Pharmacol.*, 13, 16-26); biodegradable polymers, such as poly (DL-lactide-co-glycolide) microspheres for sustained release delivery after implantation (Emerich, DF *et al.*, 1999, *Cell Transplant.*, 8, 47-58) Alkermes, Inc. Cambridge, MA; and loaded nanoparticles, such as those made of polybutylcyanoacrylate, which can  
15 deliver drugs across the blood brain barrier and can alter neuronal uptake mechanisms (*Prog Neuropsychopharmacol Biol Psychiatry*, 23, 941-949, 1999). Other non-limiting examples of delivery strategies, including CNS delivery of the nucleic acid molecules of the instant invention include material described in Boado *et al.*, 1998, *J. Pharm. Sci.*, 87, 1308-1315; Tyler *et al.*, 1999, *FEBS Lett.*, 421, 280-284; Pardridge *et al.*, 1995, *PNAS USA.*, 92, 5592-  
20 5596; Boado, 1995, *Adv. Drug Delivery Rev.*, 15, 73-107; Aldrian-Herrada *et al.*, 1998, *Nucleic Acids Res.*, 26, 4910-4916; and Tyler *et al.*, 1999, *PNAS USA.*, 96, 7053-7058. All  
25 these references are hereby incorporated herein by reference.

The invention also features the use of the composition comprising surface-modified liposomes containing poly (ethylene glycol) lipids (PEG-modified, or long-circulating liposomes or stealth liposomes). Nucleic acid molecules of the invention can also comprise covalently attached PEG molecules of various molecular weights. These formulations offer a method for increasing the accumulation of drugs in target tissues. This class of drug carriers resists opsonization and elimination by the mononuclear phagocytic system (MPS or RES), thereby enabling longer blood circulation times and enhanced tissue exposure for the  
30 35 encapsulated drug (Lasic *et al.* *Chem. Rev.* 1995, 95, 2601-2627; Ishiwata *et al.*, *Chem.*

*Pharm. Bull.* 1995, 43, 1005-1011). Such liposomes have been shown to accumulate selectively in tumors, presumably by extravasation and capture in the neovascularized target tissues (Lasic *et al.*, *Science* 1995, 267, 1275-1276; Oku *et al.*, 1995, *Biochim. Biophys. Acta*, 1238, 86-90). The long-circulating liposomes enhance the pharmacokinetics and 5 pharmacodynamics of DNA and RNA, particularly compared to conventional cationic liposomes which are known to accumulate in tissues of the MPS (Liu *et al.*, *J. Biol. Chem.* 1995, 42, 24864-24870; Choi *et al.*, International PCT Publication No. WO 96/10391; Ansell *et al.*, International PCT Publication No. WO 96/10390; Holland *et al.*, International PCT Publication No. WO 96/10392; all of which are incorporated by reference herein). Long- 10 circulating liposomes are also likely to protect drugs from nuclease degradation to a greater extent compared to cationic liposomes, based on their ability to avoid accumulation in metabolically aggressive MPS tissues such as the liver and spleen. All of these references are incorporated by reference herein.

The present invention also includes compositions prepared for storage or administration 15 which include a pharmaceutically effective amount of the desired compounds in a pharmaceutically acceptable carrier or diluent. Acceptable carriers or diluents for therapeutic use are well known in the pharmaceutical art, and are described, for example, in *Remington's Pharmaceutical Sciences*, Mack Publishing Co. (A.R. Gennaro edit. 1985) hereby incorporated by reference herein. For example, preservatives, stabilizers, dyes and flavoring 20 agents can be provided. These include sodium benzoate, sorbic acid and esters of *p*-hydroxybenzoic acid. In addition, antioxidants and suspending agents can be used.

A pharmaceutically effective dose is that dose required to prevent, inhibit the occurrence, or treat (alleviate a symptom to some extent, preferably all of the symptoms) of a disease state. The pharmaceutically effective dose depends on the type of disease, the 25 composition used, the route of administration, the type of mammal being treated, the physical characteristics of the specific mammal under consideration, concurrent medication, and other factors which those skilled in the medical arts will recognize. Generally, an amount between 0.1 mg/kg and 100 mg/kg body weight/day of active ingredients is administered dependent upon potency of the negatively charged polymer.

30 The nucleic acid molecules of the invention and formulations thereof can be administered orally, topically, parenterally, by inhalation or spray or rectally in dosage unit formulations containing conventional non-toxic pharmaceutically acceptable carriers, adjuvants and vehicles. The term parenteral as used herein includes percutaneous, subcutaneous, intravascular (e.g., intravenous), intramuscular, or intrathecal injection or

infusion techniques and the like. In addition, there is provided a pharmaceutical formulation comprising a nucleic acid molecule of the invention and a pharmaceutically acceptable carrier. One or more nucleic acid molecules of the invention can be present in association with one or more non-toxic pharmaceutically acceptable carriers and/or diluents and/or 5 adjuvants, and if desired other active ingredients. The pharmaceutical compositions containing nucleic acid molecules of the invention can be in a form suitable for oral use, for example, as tablets, troches, lozenges, aqueous or oily suspensions, dispersible powders or granules, emulsion, hard or soft capsules, or syrups or elixirs.

Compositions intended for oral use can be prepared according to any method known to 10 the art for the manufacture of pharmaceutical compositions and such compositions can contain one or more such sweetening agents, flavoring agents, coloring agents or preservative agents in order to provide pharmaceutically elegant and palatable preparations. Tablets contain the active ingredient in admixture with non-toxic pharmaceutically acceptable excipients that are suitable for the manufacture of tablets. These excipients can be for 15 example, inert diluents, such as calcium carbonate, sodium carbonate, lactose, calcium phosphate or sodium phosphate; granulating and disintegrating agents, for example, corn starch, or alginic acid; binding agents, for example starch, gelatin or acacia, and lubricating agents, for example magnesium stearate, stearic acid or talc. The tablets can be uncoated or they can be coated by known techniques. In some cases such coatings can be prepared by 20 known techniques to delay disintegration and absorption in the gastrointestinal tract and thereby provide a sustained action over a longer period. For example, a time delay material such as glycetyl monostearate or glycetyl distearate can be employed.

Formulations for oral use can also be presented as hard gelatin capsules wherein the active ingredient is mixed with an inert solid diluent, for example, calcium carbonate, calcium 25 phosphate or kaolin, or as soft gelatin capsules wherein the active ingredient is mixed with water or an oil medium, for example peanut oil, liquid paraffin or olive oil.

Aqueous suspensions contain the active materials in admixture with excipients suitable for the manufacture of aqueous suspensions. Such excipients are suspending agents, for example sodium carboxymethylcellulose, methylcellulose, hydropropyl-methylcellulose, 30 sodium alginate, polyvinylpyrrolidone, gum tragacanth and gum acacia; dispersing or wetting agents can be a naturally-occurring phosphatide, for example, lecithin, or condensation products of an alkylene oxide with fatty acids, for example polyoxyethylene stearate, or condensation products of ethylene oxide with long chain aliphatic alcohols, for example heptadecaethyleneoxycetanol, or condensation products of ethylene oxide with partial esters

derived from fatty acids and a hexitol such as polyoxyethylene sorbitol monooleate, or condensation products of ethylene oxide with partial esters derived from fatty acids and hexitol anhydrides, for example polyethylene sorbitan monooleate. The aqueous suspensions can also contain one or more preservatives, for example ethyl, or n-propyl p-hydroxybenzoate, one or more coloring agents, one or more flavoring agents, and one or more sweetening agents, such as sucrose or saccharin.

Oily suspensions can be formulated by suspending the active ingredients in a vegetable oil, for example arachis oil, olive oil, sesame oil or coconut oil, or in a mineral oil such as liquid paraffin. The oily suspensions can contain a thickening agent, for example beeswax, hard paraffin or cetyl alcohol. Sweetening agents and flavoring agents can be added to provide palatable oral preparations. These compositions can be preserved by the addition of an anti-oxidant such as ascorbic acid.

Dispersible powders and granules suitable for preparation of an aqueous suspension by the addition of water provide the active ingredient in admixture with a dispersing or wetting agent, suspending agent and one or more preservatives. Suitable dispersing or wetting agents or suspending agents are exemplified by those already mentioned above. Additional excipients, for example sweetening, flavoring and coloring agents, can also be present.

Pharmaceutical compositions of the invention can also be in the form of oil-in-water emulsions. The oily phase can be a vegetable oil or a mineral oil or mixtures of these. Suitable emulsifying agents can be naturally-occurring gums, for example gum acacia or gum tragacanth, naturally-occurring phosphatides, for example soy bean, lecithin, and esters or partial esters derived from fatty acids and hexitol anhydrides, for example sorbitan monooleate, and condensation products of the said partial esters with ethylene oxide, for example polyoxyethylene sorbitan monooleate. The emulsions can also contain sweetening and flavoring agents.

Syrups and elixirs can be formulated with sweetening agents, for example glycerol, propylene glycol, sorbitol, glucose or sucrose. Such formulations can also contain a demulcent, a preservative and flavoring and coloring agents. The pharmaceutical compositions can be in the form of a sterile injectable aqueous or oleaginous suspension. This suspension can be formulated according to the known art using those suitable dispersing or wetting agents and suspending agents that have been mentioned above. The sterile injectable preparation can also be a sterile injectable solution or suspension in a non-toxic parentally acceptable diluent or solvent, for example as a solution in 1,3-butanediol. Among the acceptable vehicles and solvents that can be employed are water, Ringer's solution and

isotonic sodium chloride solution. In addition, sterile, fixed oils are conventionally employed as a solvent or suspending medium. For this purpose any bland fixed oil can be employed including synthetic mono-or diglycerides. In addition, fatty acids such as oleic acid find use in the preparation of injectables.

5       The nucleic acid molecules of the invention can also be administered in the form of suppositories, e.g., for rectal administration of the drug. These compositions can be prepared by mixing the drug with a suitable non-irritating excipient that is solid at ordinary temperatures but liquid at the rectal temperature and will therefore melt in the rectum to release the drug. Such materials include cocoa butter and polyethylene glycols.

10      Nucleic acid molecules of the invention can be administered parenterally in a sterile medium. The drug, depending on the vehicle and concentration used, can either be suspended or dissolved in the vehicle. Advantageously, adjuvants such as local anesthetics, preservatives and buffering agents can be dissolved in the vehicle.

15      Dosage levels of the order of from about 0.1 mg to about 140 mg per kilogram of body weight per day are useful in the treatment of the above-indicated conditions (about 0.5 mg to about 7 g per patient per day). The amount of active ingredient that can be combined with the carrier materials to produce a single dosage form varies depending upon the host treated and the particular mode of administration. Dosage unit forms generally contain between from about 1 mg to about 500 mg of an active ingredient.

20      It is understood that the specific dose level for any particular patient depends upon a variety of factors including the activity of the specific compound employed, the age, body weight, general health, sex, diet, time of administration, route of administration, and rate of excretion, drug combination and the severity of the particular disease undergoing therapy.

25      For administration to non-human animals, the composition can also be added to the animal feed or drinking water. It can be convenient to formulate the animal feed and drinking water compositions so that the animal takes in a therapeutically appropriate quantity of the composition along with its diet. It can also be convenient to present the composition as a premix for addition to the feed or drinking water.

30      The nucleic acid molecules of the present invention can also be administered to a patient in combination with other therapeutic compounds to increase the overall therapeutic effect. The use of multiple compounds to treat an indication can increase the beneficial effects while reducing the presence of side effects.

Alternatively, certain of the nucleic acid molecules of the instant invention can be expressed within cells from eukaryotic promoters (e.g., Izant and Weintraub, 1985, *Science*, 229, 345; McGarry and Lindquist, 1986, *Proc. Natl. Acad. Sci., USA* 83, 399; Scanlon *et al.*, 1991, *Proc. Natl. Acad. Sci. USA*, 88, 10591-5; Kashani-Sabet *et al.*, 1992, *Antisense Res. Dev.*, 2, 3-15; Dropulic *et al.*, 1992, *J. Virol.*, 66, 1432-41; Weerasinghe *et al.*, 1991, *J. Virol.*, 65, 5531-4; Ojwang *et al.*, 1992, *Proc. Natl. Acad. Sci. USA*, 89, 10802-6; Chen *et al.*, 1992, *Nucleic Acids Res.*, 20, 4581-9; Sarver *et al.*, 1990 *Science*, 247, 1222-1225; Thompson *et al.*, 1995, *Nucleic Acids Res.*, 23, 2259; Good *et al.*, 1997, *Gene Therapy*, 4, 45; all of these references are hereby incorporated in their totalities by reference herein). Those skilled in the art realize that any nucleic acid can be expressed in eukaryotic cells from the appropriate DNA/RNA vector. The activity of such nucleic acids can be augmented by their release from the primary transcript by a enzymatic nucleic acid (Draper *et al.*, PCT WO 93/23569, and Sullivan *et al.*, PCT WO 94/02595; Ohkawa *et al.*, 1992, *Nucleic Acids Symp. Ser.*, 27, 15-6; Taira *et al.*, 1991, *Nucleic Acids Res.*, 19, 5125-30; Ventura *et al.*, 1993, *Nucleic Acids Res.*, 21, 3249-55; Chowrira *et al.*, 1994, *J. Biol. Chem.*, 269, 25856; all of these references are hereby incorporated in their totalities by reference herein). Gene therapy approaches specific to the CNS are described by Blesch *et al.*, 2000, *Drug News Perspect.*, 13, 269-280; Peterson *et al.*, 2000, *Cent. Nerv. Syst. Dis.*, 485-508; Peel and Klein, 2000, *J. Neurosci. Methods*, 98, 95-104; Hagihara *et al.*, 2000, *Gene Ther.*, 7, 759-763; and Herrlinger *et al.*, 2000, *Methods Mol. Med.*, 35, 287-312. AAV-mediated delivery of nucleic acid to cells of the nervous system is further described by Kaplitt *et al.*, US 6,180,613.

In another aspect of the invention, RNA molecules of the present invention are preferably expressed from transcription units (see for example Couture *et al.*, 1996, *TIG.*, 12, 510) inserted into DNA or RNA vectors. The recombinant vectors are preferably DNA plasmids or viral vectors. Ribozyme expressing viral vectors can be constructed based on, but not limited to, adeno-associated virus, retrovirus, adenovirus, or alphavirus. Preferably, the recombinant vectors capable of expressing the nucleic acid molecules are delivered as described above, and persist in target cells. Alternatively, viral vectors can be used that provide for transient expression of nucleic acid molecules. Such vectors can be repeatedly administered as necessary. Once expressed, the nucleic acid molecule binds to the target mRNA. Delivery of nucleic acid molecule expressing vectors can be systemic, such as by intravenous or intra-muscular administration, by administration to target cells ex-planted from the patient followed by reintroduction into the patient, or by any other means that would allow for introduction into the desired target cell (for a review see Couture *et al.*, 1996, *TIG.*, 12, 510).

In one aspect the invention features an expression vector comprising a nucleic acid sequence encoding at least one of the nucleic acid molecules of the instant invention. The nucleic acid sequence encoding the nucleic acid molecule of the instant invention is operably linked in a manner which allows expression of that nucleic acid molecule.

- 5        In another aspect the invention features an expression vector comprising: a) a transcription initiation region (e.g., eukaryotic pol I, II or III initiation region); b) a transcription termination region (e.g., eukaryotic pol I, II or III termination region); c) a nucleic acid sequence encoding at least one of the nucleic acid catalyst of the instant invention; and wherein said sequence is operably linked to said initiation region and said  
10      10 termination region, in a manner which allows expression and/or delivery of said nucleic acid molecule. The vector can optionally include an open reading frame (ORF) for a protein operably linked on the 5' side or the 3'-side of the sequence encoding the nucleic acid catalyst of the invention; and/or an intron (intervening sequences).

Transcription of the nucleic acid molecule sequences are driven from a promoter for  
15      15 eukaryotic RNA polymerase I (pol I), RNA polymerase II (pol II), or RNA polymerase III (pol III). Transcripts from pol II or pol III promoters are expressed at high levels in all cells; the levels of a given pol II promoter in a given cell type depends on the nature of the gene regulatory sequences (enhancers, silencers, etc.) present nearby. Prokaryotic RNA polymerase promoters are also used, providing that the prokaryotic RNA polymerase enzyme  
20      20 is expressed in the appropriate cells (Elroy-Stein and Moss, 1990, *Proc. Natl. Acad. Sci. U S A*, 87, 6743-7; Gao and Huang 1993, *Nucleic Acids Res.*, 21, 2867-72; Lieber *et al.*, 1993, *Methods Enzymol.*, 217, 47-66; Zhou *et al.*, 1990, *Mol. Cell. Biol.*, 10, 4529-37). All of these references are incorporated by reference herein. Several investigators have demonstrated that nucleic acid molecules, such as ribozymes expressed from such promoters  
25      25 can function in mammalian cells (e.g. Kashani-Sabet *et al.*, 1992, *Antisense Res. Dev.*, 2, 3-15; Ojwang *et al.* 1992, *Proc. Natl. Acad. Sci. U S A*, 89, 10802-6; Chen *et al.*, 1992, *Nucleic Acids Res.*, 20, 4581-9; Yu *et al.*, 1993, *Proc. Natl. Acad. Sci. U S A*, 90, 6340-4; L'Huillier *et al.*, 1992, *EMBO J.*, 11, 4411-8; Lisziewicz *et al.*, 1993, *Proc. Natl. Acad. Sci. U. S. A*, 90, 8000-4; Thompson *et al.*, 1995, *Nucleic Acids Res.*, 23, 2259; Sullenger & Cech,  
30      30 1993, *Science*, 262, 1566). More specifically, transcription units such as the ones derived from genes encoding U6 small nuclear (snRNA), transfer RNA (tRNA) and adenovirus VA RNA are useful in generating high concentrations of desired RNA molecules such as ribozymes in cells (Thompson *et al.*, *supra*; Couture and Stinchcomb, 1996, *supra*; Noonberg *et al.*, 1994, *Nucleic Acid Res.*, 22, 2830; Noonberg *et al.*, US Patent No. 5,624,803; Good *et al.*, 1997, *Gene Ther.*, 4, 45; Beigelman *et al.*, International PCT Publication No. WO  
35      35

- 96/18736; all of these publications are incorporated by reference herein. The above ribozyme transcription units can be incorporated into a variety of vectors for introduction into mammalian cells, including but not restricted to, plasmid DNA vectors, viral DNA vectors (such as adenovirus or adeno-associated virus vectors), or viral RNA vectors (such as 5 retroviral or alphavirus vectors) (for a review see Couture and Stinchcomb, 1996, *supra*).

In another aspect the invention features an expression vector comprising nucleic acid sequence encoding at least one of the nucleic acid molecules of the invention, in a manner which allows expression of that nucleic acid molecule. The expression vector comprises in one embodiment; a) a transcription initiation region; b) a transcription termination region; c) 10 a nucleic acid sequence encoding at least one said nucleic acid molecule; and wherein said sequence is operably linked to said initiation region and said termination region, in a manner which allows expression and/or delivery of said nucleic acid molecule.

In another embodiment the expression vector comprises: a) a transcription initiation region; b) a transcription termination region; c) an open reading frame; d) a nucleic acid 15 sequence encoding at least one said nucleic acid molecule, wherein said sequence is operably linked to the 3'-end of said open reading frame; and wherein said sequence is operably linked to said initiation region, said open reading frame and said termination region, in a manner which allows expression and/or delivery of said nucleic acid molecule. In yet another embodiment the expression vector comprises: a) a transcription initiation region; b) a transcription termination region; c) an intron; d) a nucleic acid sequence encoding at least one 20 said nucleic acid molecule; and wherein said sequence is operably linked to said initiation region, said intron and said termination region, in a manner which allows expression and/or delivery of said nucleic acid molecule.

In another embodiment, the expression vector comprises: a) a transcription initiation region; b) a transcription termination region; c) an intron; d) an open reading frame; e) a 25 nucleic acid sequence encoding at least one said nucleic acid molecule, wherein said sequence is operably linked to the 3'-end of said open reading frame; and wherein said sequence is operably linked to said initiation region, said intron, said open reading frame and said termination region, in a manner which allows expression and/or delivery of said nucleic acid 30 molecule.

Flt-1 (VEGFR1), KDR (VEGFR2) and/or flk-1 are attractive nucleic acid-based therapeutic targets by several criteria. The interaction between VEGF and VEGF-R is well-established. Efficacy can be tested in well-defined and predictive animal models. Finally, the disease conditions are serious and current therapies are inadequate. Whereas protein-based

therapies are designed to affect VEGF activity, nucleic acid-based therapy based on the molecules and methods described herein provides a direct and elegant approach to directly modulate flt-1, KDR and/or flk-1 expression.

Because VEGFR1 and VEGFR2 mRNAs are highly homologous in certain regions, 5 some nucleic acid target sites are also homologous. In this case, a single nucleic acid molecule of the invention can target both VEGFR1 and VEGFR2 mRNAs. At partially homologous sites, a single nucleic acid molecule can sometimes be designed to accommodate a site on both mRNAs by including G/U base pairing. For example, if there is a G present in a enzymatic nucleic acid target site in VEGFR1 mRNA at the same position there is an A in 10 the VEGFR2 enzymatic nucleic acid target site, the enzymatic nucleic acid can be synthesized with a U at the complementary position and it will bind both to sites. The advantage of one enzymatic nucleic acid that targets both VEGFR1 and VEGFR2 mRNAs is clear, especially in cases where both VEGF receptors may contribute to the progression of angiogenesis in the disease state.

15

### Examples

The following are non-limiting examples showing the selection, isolation, synthesis and activity of exemplary nucleic acids of the instant invention.

20 The following examples demonstrate the selection and design of antisense, aptamer, dsRNA, allozyme, hammerhead, DNAzyme, NCH, Amberzyme, Zinzyme, or G-Cleaver ribozyme molecules and binding/cleavage sites within VEGF, VEGFR1 and/or VEGFR2 RNA.

#### Example 1: Enzymatic nucleic acid-mediated inhibition of angiogenesis *in vivo*

25 The study described below was performed to assess the anti-angiogenic activity of hammerhead ribozymes targeted against flt-1 4229 site (SED ID NO: 5977) in the rat cornea model of VEGF induced angiogenesis (see above). These ribozymes have either active or inactive catalytic core and either bind and cleave or just bind to VEGF-R mRNA of the flt-1 subtype. The active ribozymes, that are able to bind and cleave the target RNA, have been shown to inhibit (<sup>125</sup>I-labeled) VEGF binding in cultured endothelial cells and produce a dose-dependent decrease in VEGF induced endothelial cell proliferation in these cells. The 30 catalytically inactive forms of these ribozymes, which can only bind to the RNA but cannot catalyze RNA cleavage, failed to inhibit VEGF binding and failed to decrease VEGF induced endothelial cell proliferation. The ribozymes and VEGF were co-delivered using the filter

disk method: Nitrocellulose filter disks (Millipore<sup>®</sup>) of 0.057 diameter were immersed in appropriate solutions and were surgically implanted in rat cornea as described by Pandey *et al.*, *supra*. This delivery method has been shown to deliver rhodamine-labeled free ribozyme to scleral cells and, in all likelihood cells of the pericorneal vascular plexus. Since the active 5 ribozymes show cell culture efficacy and can be delivered to the target site using the disk method, it is essential that these ribozymes be assessed for *in vivo* anti-angiogenic activity.

The stimulus for angiogenesis in this study was the treatment of the filter disk with 30  $\mu\text{M}$  VEGF which is implanted within the cornea's stroma. This dose yields reproducible neovascularization stemming from the pericorneal vascular plexus growing toward the disk in 10 a dose-response study 5 days following implant. Filter disks treated only with the vehicle for VEGF show no angiogenic response. The ribozymes were co-administered with VEGF on a disk in two different ribozyme concentrations. One concern with the simultaneous administration is that the ribozymes will not be able to inhibit angiogenesis since VEGF receptors can be stimulated. However, we have observed that in low VEGF doses, the 15 neovascular response reverts to normal suggesting that the VEGF stimulus is essential for maintaining the angiogenic response. Blocking the production of VEGF receptors using simultaneous administration of anti-VEGF-R mRNA ribozymes could attenuate the normal neovascularization induced by the filter disk treated with VEGF.

Materials and Methods:

20 1. Stock hammerhead ribozyme solutions:

a. flt-1 4229 (786  $\mu\text{M}$ )— Active

b. flt-1 4229 (736  $\mu\text{M}$ )— Inactive

2. Experimental solutions/groups:

Group 1 Solution 1 Control VEGF solution: 30  $\mu\text{M}$  in 82mM Tris base

25 Group 2 Solution 2 flt-1 4229 (1  $\mu\text{g}/\mu\text{L}$ ) in 30  $\mu\text{M}$  VEGF/82 mM Tris base

Group 3 Solution 3 flt-1 4229 (10  $\mu\text{g}/\mu\text{L}$ ) in 30  $\mu\text{M}$  VEGF/82 mM Tris base

Group 4 Solution 4 No VEGF, flt-1 4229 (10  $\mu\text{g}/\mu\text{L}$ ) in 82 mM Tris base

Group 5 Solution 5 No VEGF, No ribozyme in 82 mM Tris base

10 eyes per group, 5 animals (Since they have similar molecular weights, the molar concentrations should be essentially similar).

Each solution (VEGF and RIBOZYMES) were prepared as a 2X solution for 1:1 mixing for final concentrations above, with the exception of solution 1 in which VEGF was 2X and  
5 diluted with ribozyme diluent (sterile water).

### 3. VEGF Solutions

The 2X VEGF solution (60  $\mu$ M) was prepared from a stock of 0.82  $\mu$ g/ $\mu$ L in 50 mM Tris base. 200  $\mu$ L of VEGF stock was concentrated by speed vac to a final volume of 60.8  $\mu$ L, for a final concentration of 2.7  $\mu$ g/ $\mu$ L or 60  $\mu$ M. Six 10  $\mu$ L aliquots was prepared for daily  
10 mixing. 2X solutions for VEGF and Ribozyme was stored at 4°C until the day of the surgery. Solutions were mixed for each day of surgery. Original 2X solutions was prepared on the day before the first day of the surgery.

### 4. Surgical Solutions:

#### Anesthesia:

15 stock ketamine hydrochloride 100 mg/mL

stock xylazine hydrochloride 20 mg/mL

stock acepromazine 10 mg/mL

Final anesthesia solution: 50 mg/mL ketamine, 10 mg/mL xylazine, and 0.5 mg/mL acepromazine

20 5% povidone iodine for ophthalmic surgical wash

2% lidocaine (sterile) for ophthalmic administration (2 drops per eye)

sterile 0.9% NaCl for ophthalmic irrigation

### 5. Surgical Methods:

Standard surgical procedure as described in Pandey *et al.*, *supra*. Filter disks were  
25 incubated in 1  $\mu$ L of each solution for approximately 30 minutes prior to implantation.

### 6. Experimental Protocol:

The animal cornea were treated with the treatment groups as described above. Animals were allowed to recover for 5 days after treatment with daily observation (scoring 0 - 3). On the fifth day animals were euthanized and digital images of each eye was obtained for quantitaion using Image Pro Plus. Quantitated neovascular surface area were analyzed by 5 ANOVA followed by two post-hoc tests including Dunnets and Tukey-Kramer tests for significance at the 95% confidence level. Dunnets provide information on the significance between the differences within the means of treatments vs. controls while Tukey-Kramer provide information on the significance of differences within the means of each group.

10 The *flt-1* 4229 (SEQ ID NO: 5977) active hammerhead ribozyme at both concentrations was effective at inhibiting angiogenesis while the inactive ribozyme did not show any significant reduction in angiogenesis. A statistically significant reduction in neovascular surface area was observed only with active ribozymes. This result clearly shows that the ribozymes are capable of significantly inhibiting angiogenesis *in vivo*. Specifically, given ribozyme mechanism of action, the observed inhibition is by the binding and cleavage 15 of target RNA by ribozymes.

Example 2: Bioactivity of anti-angiogenesis ribozymes targeting *flt-1* and *kdr* RNA

#### MATERIALS AND METHODS

20 **Ribozymes :** Hammerhead ribozymes and controls designed to have attenuated activity (attenuated controls) were synthesized and purified as previously described above. The attenuated ribozyme controls maintain the binding arm sequence of the parent ribozyme and thus are still capable of binding to the mRNA target. However, they have two nucleotide changes in the core sequence that substantially reduce their ability to carry out the cleavage reaction. Ribozymes were designed to target *Flt-1* or *KDR* mRNA sites conserved in human, mouse, and rat. In general, ribozymes with binding arms of seven nucleotides were designed 25 and tested. If, however, only six nucleotides surrounding the cleavage site were conserved in all three species, six nucleotide binding arms were used. Data are presented herein for 2'-NH<sub>2</sub> uridine modified ribozymes in cell proliferation studies and for 2'-C-allyl uridine modified ribozymes in RNase protection, *in vitro* cleavage and corneal studies.

30 ***In vitro* ribozyme cleavage assays:** *In vitro* RNA cleavage rates on a 15 nucleotide synthetic RNA substrate were measured as previously described above.

**Cell culture:** Human dermal microvascular endothelial cells (HMVEC-d, Clonetics Corp.) were maintained at 37°C in flasks or plates coated with 1.5% porcine skin gelatin (300

bloom, Sigma) in Growth medium (Clonetics Corp.) supplemented with 10-20% fetal bovine serum (FBS, Hyclone). Cells were grown to confluence and used up to the seventh passage. Stimulation medium consisted of 50% Sigma 99 media and 50% RPMI 1640 with L-glutamine and additional supplementation with 10 µg/mL Insulin-Transferrin-Selenium (Gibco BRL) and 10% FBS. Cell growth was stimulated by incubation in Stimulation medium supplemented with 20 ng/mL of either VEGF<sub>165</sub> or bFGF. VEGF<sub>165</sub> (165 amino acids) was selected for cell culture and animal studies because it is the predominant form of the four native forms of VEGF generated by alternative mRNA splicing. Cell culture assays were carried out in triplicate.

10       **Ribozyme and ribozyme/LIPOFECTAMINE™ formulations:**

15       *Cell culture:* Ribozymes or attenuated controls (50-200 nM) were formulated for cell culture studies and used immediately. Formulations were carried out with LIPOFECTAMINE™ (Gibco BRL) at a 3:1 lipid to phosphate charge ratio in serum-free medium (OPTI-MEM™, Gibco BRL) by mixing for 20 minutes at room temperature. For example, a 3:1 lipid to phosphate charge ratio was established by complexing 200 nM ribozyme with 10.8 µg/µL LIPOFECTAMINE™ (13.5 µM DOSPA).

20       *In vivo:* For corneal studies, lyophilized ribozyme or attenuated controls were resuspended in sterile water at a final stock concentration of 170 µg/µL (highest dose). Lower doses (1.7-50 µg/µL) were prepared by serial dilution in sterile water.

25       **Proliferation assay:** HMVEC-d were seeded ( $5 \times 10^3$  cells/well) in 48-well plates (Costar) and incubated 24-30 hours in Growth medium at 37°C. After removal of the Growth medium, cells were treated with 50-200 nM LIPOFECTAMINE™ complexes of ribozyme or attenuated controls for 2 hours in OPTI-MEM™. The ribozyme/control-containing medium was removed and the cells were washed extensively in 1X PBS. The medium was then replaced with Stimulation medium or Stimulation medium supplemented with 20 ng/mL VEGF<sub>165</sub> or bFGF. After 48 hours, the cell number was determined using a Coulter™ cell counter. Data are presented as cell number per well following 48 hours of VEGF stimulation.

30       **RNAse protection assay:** HMVEC-d were seeded ( $2 \times 10^5$  cells/well) in 6-well plates (Costar) and allowed to grow 32-36 hours in Growth medium at 37°C. Cells were treated with LIPOFECTAMINE™ complexes containing 200 nM ribozyme or attenuated control for 2 h as described under "Proliferation Assay" and then incubated in Growth medium containing 20 ng/mL VEGF<sub>165</sub> for 24 hours. Cells were harvested and an RNAse protection assay was carried out using the Ambion Direct Protect kit and protocol with the exception that 50 mM

EDTA was added to the lysis buffer to eliminate the possibility of ribozyme cleavage during sample preparation. Antisense RNA probes targeting portions of *Flt-1* and *KDR* were prepared by transcription in the presence of [<sup>32</sup>P]-UTP. Samples were analyzed on polyacrylamide gels and the level of protected RNA fragments was quantified using a  
5 Molecular Dynamics PhosphorImager. The levels of *Flt-1* and *KDR* were normalized to the level of cyclophilin (human cyclophilin probe template, Ambion) in each sample. The coefficient of variation for cyclophilin levels was 11% [265940 cpm ± 29386 (SD)] for all conditions tested here (*i.e.* in the presence of either active ribozymes or attenuated controls). Thus, cyclophilin is useful as an internal standard in these studies.

10       **Rat corneal pocket assay of VEGF-induced angiogenesis:**

*Animal guidelines and anesthesia.* Animal housing and experimentation adhered to standards outlined in the 1996 Guide for the Care and Use of Laboratory Animals (National Research Council). Male Sprague Dawley rats (250-300 g) were anesthetized with ketamine (50 mg/kg), xylazine (10 mg/kg), and acepromazine (0.5 mg/kg) administered intramuscularly  
15 (im). The level of anesthesia was monitored every 2-3 min by applying hind limb paw pressure and examining for limb withdrawal. Atropine (0.4 mg/kg, im) was also administered to prevent potential corneal reflex-induced bradycardia.

*Preparation of VEGF soaked disk.* For corneal implantation, 0.57 mm diameter nitrocellulose disks, prepared from 0.45 µm pore diameter nitrocellulose filter membranes  
20 (Millipore Corporation), were soaked for 30 min in 1 µL of 30 µM VEGF<sub>165</sub> in 82 mM Tris-HCl (pH 6.9) in covered petri dishes on ice.

*Corneal surgery.* The rat corneal model used in this study was a modified from Koch *et al.* *Supra* and Pandey *et al.*, *supra*. Briefly, corneas were irrigated with 0.5% povidone iodine solution followed by normal saline and two drops of 2% lidocaine. Under a dissecting microscope (Leica MZ-6), a stromal pocket was created and a presoaked filter disk (see above) was inserted into the pocket such that its edge was 1 mm from the corneal limbus.  
25

*Intraconjunctival injection of test solutions.* Immediately after disk insertion, the tip of a 40-50 µm OD injector (constructed in our laboratory) was inserted within the conjunctival tissue 1 mm away from the edge of the corneal limbus that was directly adjacent to the  
30 VEGF-soaked filter disk. Six hundred nanoliters of test solution (ribozyme, attenuated control or sterile water vehicle) were dispensed at a rate of 1.2 µL/min using a syringe pump (KD Scientific). The injector was then removed, serially rinsed in 70% ethanol and sterile water and immersed in sterile water between each injection. Once the test solution was injected,

closure of the eyelid was maintained using microaneurism clips until the animal began to recover gross motor activity. Following treatment, animals were warmed on a heating pad at 37°C.

- Animal treatment groups/experimental protocol.* Ribozymes targeting *Flt-1* site 4229 (SEQ ID NO: 5977) and *KDR* mRNA site 726 (SEQ ID NO: 5978) were tested in the corneal model along with their attenuated controls. Five treatment groups were assigned to examine the effects of five doses of each test substance over a dose range of 1-100 µg on VEGF-stimulated angiogenesis. Negative (30 µM VEGF soaked filter disk and intraconjunctival injection of 600 nL sterile water) and no stimulus (Tris-soaked filter disk and intraconjunctival injection of sterile water) control groups were also included. Each group consisted of five animals (10 eyes) receiving the same treatment.

- Quantitation of angiogenic response.* Five days after disk implantation, animals were euthanized following im administration of 0.4 mg/kg atropine and corneas were digitally imaged. The neovascular surface area (NSA, expressed in pixels) was measured *postmortem* from blood-filled corneal vessels using computerized morphometry (Image Pro Plus, Media Cybernetics, v2.0). The individual mean NSA was determined in triplicate from three regions of identical size in the area of maximal neovascularization between the filter disk and the limbus. The number of pixels corresponding to the blood-filled corneal vessels in these regions was summated to produce an index of NSA. A group mean NSA was then calculated. Data from each treatment group were normalized to VEGF/ribozyme vehicle-treated control NSA and finally expressed as percent inhibition of VEGF-induced angiogenesis.

- Statistics.* After determining the normality of treatment group means, group mean percent inhibition of VEGF-induced angiogenesis was subjected to a one-way analysis of variance. This was followed by two post-hoc tests for significance including Dunnett's (comparison to VEGF control) and Tukey-Kramer (all other group mean comparisons) at alpha = 0.05. Statistical analyses were performed using JMP v.3.1.6 (SAS Institute).

## RESULTS

- Ribozyme-mediated reduction of VEGF-induced cell proliferation:** Ribozyme cleavage of *Flt-1* or *KDR* mRNA should result in a decrease in the density of cell surface VEGF receptors. This decrease should limit VEGF binding and consequently interfere with the mitogenic signaling induced by VEGF. To determine if cell proliferation was impacted by anti-*Flt-1* and/or anti-*KDR* ribozyme treatment, proliferation assays using cultured human microvascular cells were carried out. Ribozymes included in the proliferation assays were

initially chosen by their ability to decrease the level of VEGF binding to treated cells. In these initial studies, ribozymes targeting 20 sites in the coding region of each mRNA were screened. The most effective ribozymes against two sites in each target, *Flt-1* sites 1358 and 4229 and *KDR* sites 726 and 3950, were included in the proliferation assays reported here. In 5 addition, attenuated analogs of each ribozyme were used as controls. These attenuated controls are still capable of binding to the mRNA target since the binding arm sequence is maintained. However, these controls have two nucleotide changes in the core sequence that substantially reduce their ability to carry out the cleavage reaction.

The active ribozymes tested decreased the relative proliferation of HMVEC-d after 10 VEGF stimulation, an effect that increased with ribozyme concentration. This concentration dependency was not observed following treatment with the attenuated controls designed for these sites. In fact, little or no change in cell growth was noted following treatment with the attenuated controls, even though these controls can still bind to the specific target sequences. At 200 nM, there was a distinct "window" between the anti-proliferative effects of each 15 ribozyme and its attenuated control; a trend also observed at lower doses. This window of inhibition of proliferation (56-77% based on total cells/well) reflects the contribution of ribozyme-mediated activity. In comparison, no effect of anti-*Flt-1* or anti-*KDR* ribozymes was noted on bFGF-stimulated cell proliferation. Moreover, an irrelevant, but active, 20 ribozyme whose binding sequence is not found in either *Flt-1* or *KDR* mRNA had no effect in this assay. These data are consistent with the basic ribozyme mechanism in which binding and cleavage are necessary components. Although the relative surface distribution of *Flt-1* and *KDR* receptors in this cell type is not known, the antiproliferative effects of these ribozymes indicate that, at least in cell culture, both receptors are functionally coupled to proliferation.

25       **Specific reduction of *Flt-1* or *KDR* mRNA by ribozyme treatment:** To confirm that anti-*Flt-1* and anti-*KDR* ribozymes reduce their respective mRNA targets, cellular levels of *Flt-1* or *KDR* were quantified using an RNase protection assay with specific *Flt-1* or *KDR* probes. For each target, one ribozyme/attenuated control pair was chosen for continued study. Exposure of HMVEC-d to active ribozyme targeting *Flt-1* site 4229 decreased *Flt-1* mRNA, 30 but not *KDR* mRNA. Likewise, treatment with the active ribozyme targeting *KDR* site 726 decreased *KDR*, but not *Flt-1* mRNA. Both ribozymes decreased the level of their respective target RNA by greater than 50%. The degree of reduction associated with the corresponding attenuated controls was not greater than 13%.

*In vitro* activity of anti-*Flt* and anti-*KDR* ribozymes.

To confirm further the necessity of an active ribozyme core, *in vitro* cleavage activities were determined for the *Flt-1* site 4229 ribozyme and the *KDR* site 726 ribozyme as well as their paired attenuated controls. The first order rate constants calculated from the time-course of short substrate cleavage for the anti-*Flt-1* ribozyme and its attenuated control were  $0.081 \pm 0.0007 \text{ min}^{-1}$  and  $0.001 \pm 6 \times 10^{-5} \text{ min}^{-1}$ , respectively. For the anti-*KDR* ribozyme and its paired control, the first order rate constants were  $0.434 \pm 0.024 \text{ min}^{-1}$  and  $0.002 \pm 1 \times 10^{-4} \text{ min}^{-1}$ , respectively. Although the attenuated controls retain a very slight level of cleavage activity under these optimized conditions, the decrease in *in vitro* cleavage activity between each active ribozyme and its paired attenuated control is about two orders of magnitude.

Thus, an active core is essential for cleavage activity *in vitro* and is also necessary for ribozyme activity in cell culture.

Ribozyme-mediated reduction of VEGF-induced angiogenesis *in vivo*. To assess whether ribozymes targeting VEGF receptor mRNA could impact the complex process of angiogenesis, prototypic anti-*Flt-1* and *KDR* ribozymes that were identified in cell culture studies were screened in a rat corneal pocket assay of VEGF-induced angiogenesis. In this assay, corneas implanted with VEGF-containing filter disks exhibited a robust neovascular response in the corneal region between the disk and the corneal limbus (from which the new vessels emerge). Disks containing a vehicle solution elicited no angiogenic response. In separate studies, intraconjunctival injections of sterile water vehicle did not affect the magnitude of the VEGF-induced angiogenic response. In addition, ribozyme injections alone did not induce angiogenesis.

The dose-related effects of anti-*Flt-1* or *KDR* ribozymes on the VEGF-induced angiogenic response were then examined. The antiangiogenic effect of the anti-*Flt-1* (site 4229) and *KDR* (site 726) ribozymes and their attenuated controls over a dose range from 1 to 100 µg, respectively was determined. For both ribozymes, the maximal antiangiogenic response (48 and 36% for anti-*Flt-1* and *KDR* ribozymes, respectively) was observed at a dose of 10 µg.

The anti-*Flt-1* ribozyme produced a significantly greater antiangiogenic response than its attenuated control at 3 and 10 µg ( $p < 0.05$ ). Its attenuated control exhibited a small but significant antiangiogenic response at doses above 10 µg compared to vehicle treated VEGF controls ( $p < 0.05$ ). At its maximum, this response was not significantly greater than that observed with the lowest dose of active anti-*Flt-1* ribozyme. The anti-*KDR* ribozyme significantly inhibited angiogenesis from 3 to 30 µg ( $p < 0.05$ ). The anti-*KDR* attenuated control had no significant effect at any dose tested.

Example 3. *In vivo* inhibition of tumor growth and metastases by VEGF-R ribozymes.

- A. Lewis Lung Carcinoma Mouse Model: Ribozymes were chemically synthesized as described above. The sequence of ANGIOZYME™ bound to its target RNA is shown in Figure 1.
- 5       The tumors in this study were derived from a cell line (LLC-HM) which gives rise to reproducible numbers of spontaneous lung metastases when propagated *in vivo*. The LLC-HM line was obtained from Dr. Michael O'Reilly, Harvard University. Tumor neovascularization in Lewis lung carcinoma has been shown to be VEGF-dependent. Tumors from mice bearing LLC-HM (selected for the highly metastatic phenotype by serial propagation) were harvested 20 days post-inoculation. A tumor brei suspension was prepared from these tumors according to standard protocols. On day 0 of the study,  $0.5 \times 10^6$  viable LLC-HM tumor cells were injected subcutaneously (sc) into the dorsum or flank of previously untreated mice (100 µL injectate). Tumors were allowed to grow for a period of 3 days prior to initiating continuous intravenous administration of saline or 30 mg/kg/d 10 ANGIOZYME™ *via* Alzet mini-pumps. One set of animals was dosed from days 3 to 17, inclusive. Tumor length and width measurements and volumes were calculated according to the formula: Volume = 0.5(length)(width)<sup>2</sup>. At post-inoculation day 25, animals were euthanized and lungs harvested. The number of lung macrometastatic nodules was counted. It should be noted that metastatic foci were quantified 8 days after the cessation of dosing. 15
- 10     Ribozyme solutions were prepared to deliver to another set of animals 100, 10, 3, or 1 mg/kg/day of ANGIOZYME™ *via* Alzet mini-pumps. A total of 10 animals per dose or saline control group were surgically implanted on the left flank with osmotic mini-pumps pre-filled with the respective test solution three days following tumor inoculation. Pumps were attached to indwelling jugular vein catheters.
- 20     Figure 2 shows the antitumor effects of ANGIOZYME™. There is a statistically significant inhibition ( $p < 0.05$ ) of primary LLC-HM tumor growth in tumors grown in the flank regions compared to saline control. ANGIOZYME™ significantly reduced ( $p < 0.05$ ) the number of lung metastatic foci in animals inoculated either in the flank regions. Figure 3 illustrates the dose-dependent anti-metastatic effect of ANGIOZYME™ compared to saline 25 control.

B. Mouse Colorectal Cancer Model. KM12L4a-16 is a human colorectal cancer cell line. On day 0 of the study,  $0.5 \times 10^6$  KM12L4a-16 cells were implanted into the spleen of nude mice. Three days after tumor inoculation, Alzet minipumps were implanted and continuous subcutaneous delivery of either saline or 12, 36 or 100 mg/kg/ day of

ANGIOZYME™ was initiated. On day 5, the spleens containing the primary tumors were removed. On day 18, the Alzet minipumps were replaced with fresh pumps so that delivery of saline or ANGIOZYME™ was continuous over a 28 day period from day 3 to day 32. Animals were euthanized on day 41 and the liver tumor burden was evaluated.

5 Following treatment with 100 mg/kg/day of ANGIOZYME™, there was a significant reduction in the incidence and median number of liver metastasis (Figure 4). In saline-treated animals, the median number of metastases was 101. However, at the high dose of ANGIOZYME™ (100 mg/kg/day), the median number of metastases was zero.

10 Example 4: Effect of ANGIOZYME™ alone or in combination with chemotherapeutic agents in the mouse Lewis Lung Carcinoma Model.

#### Methods

**Tumor inoculations.** Male C57/BL6 mice, age 6 to 8 weeks, were inoculated subcutaneously in the flank with  $5 \times 10^5$  LLC-HM cells from brei preparations made from tumors grown in mice.

15 **Ribozymes and controls.** RPI4610, also known as ANGIOZYME™ (SEQ ID NO: 5977), is an anti-*Flt-1* ribozyme that targets site 4229 in the human *Flt-1* receptor mRNA (EMBL accession no. X51602). The controls tested include RPL13141, an attenuated version of RPI4610 in which four nucleotides in the catalytic core are changed so that the cleavage activity is dramatically decreased. RPL13141, however, maintains the base composition and 20 binding arms of RPI4610 and so is still capable of binding to the target site. The second control (RPI13030) also has changes to the catalytic core (three) to inhibit cleavage activity, but in addition the sequence of the binding arms has been scrambled so that it can no longer bind to the target sequence. One nucleotide in the arm of RPI13030 is also changed to maintain the same base composition as RPL4610.

25 **Ribozyme administrations.** Ribozymes and controls were resuspended in normal saline. Administration was initiated seven days following tumor inoculation. Animals either received a daily subcutaneous injection (30 mg/kg test substance) from day 7 to day 20 or were instrumented with an Alzet osmotic minipump (12 µL/day flow rate) containing a solution of ribozyme or control. Subcutaneous infusion pumps delivered the test substances 30 (30 mg/kg/day) from day 7 to 20 (14-day pumps, 420 mg/kg total test substance) or days 7-34 (28-day pumps, 840 mg/kg total test substance). Where indicated, chemotherapeutic agents were given in combination with ribozyme treatment. Cyclophosphamide was given by intraperitoneal administration on days 7, 9 and 11 (125 mg/kg). Gemcitabine was given by

intraperitoneal administration on days 8, 11 and 14 (125 mg/kg). Untreated, uninstrumented animals were used as comparison. Five animals were included in each group.

### Results

5        The antiangiogenic ribozyme, ANGIOZYME™, was tested in a model of Lewis lung carcinoma alone and in combination with two chemotherapeutic agents. Previously (see above), 30 mg/kg/day ANGIOZYME™ alone was determined to inhibit both primary tumor growth and lung metastases in a highly metastatic variant of Lewis lung (continuous 14-day iv delivery via Alzet minipump, manuscript in preparation).

10      In this study, 30 mg/kg/day ANGIOZYME™ delivered either as a daily subcutaneous bolus injection or as a continuous infusion from an Alzet minipump resulted in a delay in tumor growth. On average, tumor growth to 500 mm<sup>3</sup> was delayed by ~7 days in animals being treated with ANGIOZYME™ compared to an untreated group. Growth of tumors in animals being treated with either of two attenuated controls was delayed by only ~ 2 days.

15      ANGIOZYME™ delivered by subcutaneous bolus was also tested in combination with either Gemcytabine or cyclophosphamide. Tumor growth delay increased by about 3 days in the presence of combination therapy with ANGIOZYME™ and Gemcytabine over the effects of either treatment alone. The combination of ANGIOZYME™ and cyclophosphamide did not 20 increase tumor growth delay over that of cyclophosphamide alone, however, suboptimal doses of cyclophosphamide were not included in this study. Neither of the attenuated controls increased the effect of the chemotherapeutic agents.

25      The effect of ANGIOZYME™ on metastases to the lung was also determined in the presence and absence of additional chemotherapeutic treatment. Macrometastases to the lungs were counted in two animals in each treatment group on day 20. In the presence of ANGIOZYME™, with or without a chemotherapeutic agent, the lung metastases were reduced to zero. Treatment with either Gemcytabine or cyclophosphamide alone (mean number of metastases 4.5 and 4, respectively) were not as effective as ANGIOZYME™ alone 30 or when used in combination with ANGIOZYME™. Neither of the attenuated controls increased the effect of the chemotherapeutic agents.

35      The effect on metastases to the lung was also determined following continuous treatment with ANGIOZYME™. At day 20, an average of ~8 macrometastases were noted in the treatment groups which had been instrumented with Alzet minipumps (either 14- or 28-day pumps). This is a decrease in metastases of ~50% from the untreated group. Since

ANGIOZYME™ delivered by a daily subcutaneous bolus resulted in zero metastases (Fig.4) in the two animals counted, it is possible that the additional burden of being instrumented with the minipump contributes to a slightly decreased response to ANGIOZYME™.

Example 5: Identification of Potential Target Sites in Human VEGFR1 and/or VEGFR2 RNA

5       The sequence of human VEGFR1 and/or VEGFR2 genes are screened for accessible sites using a computer-folding algorithm. Regions of the RNA that do not form secondary folding structures and contain potential enzymatic nucleic acid molecule and/or antisense binding/cleavage sites are identified. An exemplary sequence of an enzymatic nucleic acid molecule of the invention is shown in Formula I and/or Formula II (SEQ ID Nos: 5977 and  
10 5978, respectively). Other nucleic acid molecules and targets contemplated by the invention are described in Pavco *et al.*, US Patent Application No. 09/870,161, incorporated by reference herein in its entirety. Similarly, other nucleic acid molecules of the invention, including antisense, aptamers, dsRNA, siRNA, and/or 2,5-A chimeras, can be designed to modulate the expression of the nucleic acid targets described in Pavco *et al.*, US Patent  
15 Application No. 09/870,161.

Example 6: Selection of Enzymatic Nucleic Acid Cleavage Sites in Human VEGFR1 and/or VEGFR2 RNA

Enzymatic nucleic acid molecule target sites are chosen by analyzing sequences of human VEGFR1 receptor (for example Genbank Accession No. NM\_002019), and VEGFR2 receptor (for example Genbank Accession No. NM\_002253) genes and prioritizing the sites on the basis of folding. Enzymatic nucleic acid molecules are designed that can bind each target and are individually analyzed by computer folding (Christoffersen *et al.*, 1994 *J. Mol. Struc. Theochem*, 311, 273; Jaeger *et al.*, 1989, *Proc. Natl. Acad. Sci. USA*, 86, 7706) to assess whether the enzymatic nucleic acid molecule sequences fold into the appropriate secondary structure. Those enzymatic nucleic acid molecules with unfavorable intramolecular interactions between the binding arms and the catalytic core can be eliminated from consideration. As discussed herein, varying binding arm lengths can be chosen to optimize activity. Generally, at least 4 bases on each arm are able to bind to, or otherwise interact with, the target RNA.

30      Example 7: Chemical Synthesis and Purification of Ribozymes and Antisense for Efficient Cleavage and/or blocking of VEGFR1 and/or VEGFR2 RNA

Enzymatic nucleic acid molecules and antisense constructs are designed to anneal to various sites in the RNA message. The binding arms of the enzymatic nucleic acid molecules are complementary to the target site sequences described above, while the antisense constructs are fully complementary to the target site sequences described above. RNAi molecules (dsRNA) likewise have one strand of RNA or a portion of RNA complementarity to the target site sequence or a portion of the target site sequence. For example, complementarity within the double-strand RNAi structure is formed from two separate individual RNA strands or from self-complementary areas of a topologically closed, individual RNA strand which can be optionally circular. The nucleic acid molecules were chemically synthesized. The method of synthesis used followed the procedure for normal RNA synthesis as described above and in Usman *et al.*, (1987 J. Am. Chem. Soc., 109, 7845), Scaringe *et al.*, (1990 Nucleic Acids Res., 18, 5433) and Wincott *et al.*, *supra*, and made use of common nucleic acid protecting and coupling groups, such as dimethoxytrityl at the 5'-end, and phosphoramidites at the 3'-end. The average stepwise coupling yields were typically >98%.

Nucleic acid molecules are also synthesized from DNA templates using bacteriophage T7 RNA polymerase (Milligan and Uhlenbeck, 1989, Methods Enzymol. 180, 51). Nucleic acid molecules of the invention are purified by gel electrophoresis using general methods or are purified by high pressure liquid chromatography (HPLC; See Wincott *et al.*, *supra*; the totality of which is hereby incorporated herein by reference) and are resuspended in water. Examples of sequences of chemically synthesized enzymatic nucleic acid molecules are shown in Formula I (SEQ ID NO: 5977), Formula II (SEQ ID NO: 5978) and in Pavco *et al.*, US Patent Application No. 09/870,161.

Example 8: Enzymatic Nucleic Acid Molecule Cleavage of VEGFR1 and/or VEGFR2 RNA  
25    Target *in vitro*

Enzymatic nucleic acid molecules targeted to the human VEGFR1 and/or VEGFR2 RNA are designed and synthesized as described above. These enzymatic nucleic acid molecules can be tested for cleavage activity *in vitro*, for example, using the following procedure. The target sequences and the nucleotide location within the VEGFR1 and/or VEGFR2 RNA are described in Pavco *et al.*, US Patent Application No. 09/870,161.

*Cleavage Reactions:* Full-length or partially full-length, internally-labeled target RNA for enzymatic nucleic acid molecule cleavage assay is prepared by *in vitro* transcription in the presence of [ $\alpha$ -<sup>32</sup>P] CTP, passed over a G 50 Sephadex column by spin chromatography and used as substrate RNA without further purification. Alternately, substrates are 5'-<sup>32</sup>P-end

labeled using T4 polynucleotide kinase enzyme. Assays are performed by pre-warming a 2X concentration of purified enzymatic nucleic acid molecule in enzymatic nucleic acid molecule cleavage buffer (50 mM Tris-HCl, pH 7.5 at 37°C, 10 mM MgCl<sub>2</sub>) and the cleavage reaction was initiated by adding the 2X enzymatic nucleic acid molecule mix to an equal volume of 5 substrate RNA (maximum of 1-5 nM) that was also pre-warmed in cleavage buffer. As an initial screen, assays are carried out for 1 hour at 37°C using a final concentration of either 40 nM or 1 mM enzymatic nucleic acid molecule, *i.e.*, enzymatic nucleic acid molecule excess. The reaction is quenched by the addition of an equal volume of 95% formamide, 20 mM EDTA, 0.05% bromophenol blue and 0.05% xylene cyanol after which the sample is heated 10 to 95°C for 2 minutes, quick chilled and loaded onto a denaturing polyacrylamide gel. Substrate RNA and the specific RNA cleavage products generated by enzymatic nucleic acid molecule cleavage are visualized on an autoradiograph of the gel. The percentage of cleavage is determined by Phosphor Imager® quantitation of bands representing the intact substrate and the cleavage products.

15 **Example 9: Phase I/II Study of Repetitive Dosing of ANGIOZYME™ Targeting the VEGFR1 (FLT-1) Receptor of VEGF**

A ribozyme therapeutic agent ANGIOZYME™ (SEQ ID NO: 5977), was assessed by daily subcutaneous administration in a phase I/II trial for 31 patients with refractory solid tumors. Demographic information relating to patients enrolled in the study are shown in Table III. 20 The primary study endpoint was to determine the safety and maximum tolerated dose of ANGIOZYME™. Secondary endpoints assessed ANGIOZYME™ pharmacokinetics and clinical response. Patients were treated at the following doses: 3 patients received doses of 10 mg/m<sup>2</sup>/day, 4 patients received 30 mg/m<sup>2</sup>/day, 20 patients received 100 mg/m<sup>2</sup>/day, and 4 patients received 300 mg/m<sup>2</sup>/day. All but one patient were dosed for a minimum of 29 consecutive days with 24-hour pharmacokinetic analyses on Day 1 and 29. Clinical response was assessed monthly. Results The data from 20 patients indicated that 25 ANGIOZYME™ was well tolerated, with no systemic adverse events. Figure 5 shows the plasma concentration profile of ANGIOZYME™ after a single subcutaneous dose of 10, 30, 100, or 300 mg/m<sup>2</sup>. The pharmacokinetic parameters of ANGIOZYME™ after subcutaneous 30 bolus administration are outlined in Table IV. An MTD (maximum tolerated dose) could not be established. One patient in the 300 mg/m<sup>2</sup>/d group experienced a grade 3 injection site reaction. Patients in the other groups experienced intermittent grade 1 and grade 2 injection site reactions with erythema and induration. No systemic or laboratory toxicities were observed. Pharmacokinetic analyses demonstrated dose-dependent plasma concentrations 35 with good bioavailability (70-90%), t<sub>1/2</sub> = 209-384 min, and no accumulation after repeated

doses. To date, 17/28 (61%) of evaluable patients have had stable disease for periods of one to six months and two patients (nasopharyngeal squamous cell carcinoma and melanoma) had minor clinical responses. The patient with nasopharyngeal carcinoma demonstrated central tumor necrosis as indicated by MRI. The longest period of treatment thus far has been 8  
5 months for two patients at 100 mg/m<sup>2</sup>/d (breast, peritoneal mesothelioma).

Example 10: Down-regulation of VEGFR1 gene expression to treat gynecologic neovascularization dependent conditions

One patient in the Phase I/II trial described in Example 19 was menstruating prior to enrollment in the ANGIOZYME™ monotherapy trial. After 1-2 months on trial, the patient's menstrual cycles ceased. The patient remained on trial for approximately 11 months and did not menstruate. The patient then went off the trial for about 4 months and the menstrual cycles resumed. Re-enrollment in the ANGIOZYME™ trial resulted in the patient's menstrual cycle stopping again. This clinical observation suggests that ANGIOZYME™ is interfering with the patient's menstrual cycle, perhaps by inhibiting neovascularization of uterine tissue. This data also suggests that ANGIOZYME™ has a direct effect on the endometrial tissue or an effect on LH/FSH stimulation. These results suggest the treatment or control, using ANGIOZYME™ (SEQ ID NO: 5977) and/or other nucleic acid molecules of the instant invention, of various clinical targets and/or processes associated with female reproduction and gynecologic neovascularization, such as endometriosis, birth control,  
10 gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), menopausal dysfunction, endometrial carcinoma or other condition associated with the expression of VEGFR1 and/or VEGFR2 VEGF receptors.  
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Example 11: Down-regulation of VEGFR1 in clinical setting

Twenty-seven of the patients enrolled in the Phase I/II trial described in Example 19 had day 1 (baseline) and day 43 (six-week) serum samples assayed for VEGFR1 biomarker. VEGFR1 levels were statistically different after six weeks of ANGIOZYME treatment (Figure 9). Although statistical testing involving all 27 patients showed statistical support for effects, not all patients presented with elevated levels of VEGF-R1. Since the effects of ANGIOZYME on VEGF-R1 may only be demonstrated when sufficient levels are present at baseline, a cutoff of 100 pg/mL was chosen and changes in this VEGF-R1 were re-analyzed. Ten of the 27 patients presented with baseline VEGF-R1 levels in excess of 100 pg/mL. For this subgroup VEGF-R1 levels were lower by 3-fold, p<.001. After six weeks of treatment the average (geometric mean) of VEGF-R1 decreased for this subgroup from 419 pg/ml to  
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132pg/ml, p<.001. These results show that treatment with ANGIOZYME results in a statistically significant reduction in VEGFR1 expression.

Example 22: *In vivo* inhibition of neovascularization in an ocular animal model by VEGF-R ribozymes.

5       Summary of the Mouse Model: A mouse model of proliferative retinopathy (Aiello et al., 1995, *Proc. Natl. Acad. Sci. USA* 92: 10457-10461; Robinson et al., 1996, *Proc. Natl. Acad. Sci. USA* 93: 4851-4856; Pierce et al., 1996, *Archives of Ophthalmology* 114: 1219-1228) in which neovascularization of the mouse retina is induced by exposure of 7-day old mice to 75% oxygen followed by a return to normal room air. The initial period in high  
10      oxygen causes an obliteration of developing blood vessels in the retina. Exposure to room air five days later is perceived as hypoxia by the now underperfused retina. The result is an immediate upregulation of VEGF mRNA and VEGF protein (between 6-12 hours) followed by an extensive retinal neovascularization that peaks in ~5 days. Although this model is more representative of retinopathy of prematurity than diabetic retinopathy, it is an accepted small  
15      animal model in which to study neovascular pathophysiology of the retina. In fact, intravitreal injection of certain antisense DNA constructs targeting VEGF mRNA have been found to be antiangiogenic in this model, as were soluble VEGF receptor chimeric proteins designed to bind VEGF in the vitreous humor (Aiello et al., 1995, *Proc. Natl. Acad. Sci. USA* 92: 10457-10461; Robinson et al., 1996, *Proc. Natl. Acad. Sci. USA* 93: 4851-4856; Pierce  
20      et al., 1996, *Archives of Ophthalmology* 114: 1219-1228).

Summary of experiment: The effect of an anti-*KDR/Flik-1* ribozyme on the peak level of neovascularization was tested in the mouse model described above. As shown in Figure 10, P7 mice were removed from the hyperoxic chamber and the mice received two intraocular injections (P12 and P13) in the right eye of 10 µg RPL4731, the anti-*KDR/Flik-1* ribozyme.  
25      The left eye of each mouse was treated as a control and received intraocular injections of saline. Five days after being exposed to room air, neovascular nuclei in the retina of both eyes were counted. Data are presented in Figure 11. There was a significant decrease in retinal neovascularization (~40%) compared to the control, saline-injected eyes.

30      RPL4731 sequence and chemical composition:  
          5'-ugagcu uuU GAu Gg ga aaa gcc Gaa Aag aca aB-3' (SEQ ID NO: 5978)

where:

35      uppercase G, A = ribonucleotides  
          lowercase = 2'-OMe  
          U = 2'-C-allyl uridine

B = inverted abasic nucleotide  
S = phosphorothioate internucleotide linkage

Indications

- 5        1) Tumor angiogenesis: Angiogenesis has been shown to be necessary for tumors to grow into pathological size (Folkman, 1971, *PNAS* 76, 5217-5221; Wellstein & Czubayko, 1996, *Breast Cancer Res and Treatment* 38, 109-119). In addition, it allows tumor cells to travel through the circulatory system during metastasis. Increased levels of gene expression of a number of angiogenic factors such as vascular endothelial growth factor (VEGF) have  
10      been reported in vascularized and edema-associated brain tumors (Berkman *et al.*, 1993 *J. Clin. Invest.* 91, 153). A more direct demonstration of the role of VEGF in tumor angiogenesis was demonstrated by Jim Kim *et al.*, 1993 *Nature* 362, 841 wherein, monoclonal antibodies against VEGF were successfully used to inhibit the growth of rhabdomyosarcoma, glioblastoma multiforme cells in nude mice. Similarly, expression of a dominant negative  
15      mutated form of the flt-1 VEGF receptor inhibits vascularization induced by human glioblastoma cells in nude mice (Millauer *et al.*, 1994, *Nature* 367, 576). Specific tumor/cancer types that can be targeted using the nucleic acid molecules of the invention include but are not limited to the tumor/cancer types described under Diagnosis in Table III.
- 20        2) Ocular diseases: Neovascularization has been shown to cause or exacerbate ocular diseases including but not limited to, macular degeneration, neovascular glaucoma, diabetic retinopathy, myopic degeneration, and trachoma (Norrbj, 1997, *APMIS* 105, 417-437). Aiello *et al.*, 1994 *New Engl. J. Med.* 331, 1480, showed that the ocular fluid, of a majority of patients suffering from diabetic retinopathy and other retinal disorders, contains a high concentration of VEGF. Miller *et al.*, 1994 *Am. J. Pathol.* 145, 574, reported elevated levels  
25      of VEGF mRNA in patients suffering from retinal ischemia. These observations support a direct role for VEGF in ocular diseases. Other factors including those that stimulate VEGF synthesis may also contribute to these indications.
- 30        3) Dermatological Disorders: Many indications have been identified which may be angiogenesis dependent including but not limited to psoriasis, verruca vulgaris, angiofibroma of tuberous sclerosis, port-wine stains, Sturge Weber syndrome, Kippel-Trenaunay-Weber syndrome, and Osler-Weber-Rendu syndrome (Norrbj, *supra*). Intradermal injection of the angiogenic factor b-FGF demonstrated angiogenesis in nude mice (Weckbecker *et al.*, 1992, *Angiogenesis: Key principles-Science-Technology-Medicine*, ed R. Steiner) Detmar *et al.*, 1994 *J. Exp. Med.* 180, 1141 reported that VEGF and its receptors were over-expressed in

psoriatic skin and psoriatic dermal microvessels, suggesting that VEGF plays a significant role in psoriasis.

4) Rheumatoid arthritis: Immunohistochemistry and *in situ* hybridization studies on tissues from the joints of patients suffering from rheumatoid arthritis show an increased level 5 of VEGF and its receptors (Fava *et al.*, 1994 *J. Exp. Med.* 180, 341). Additionally, Koch *et al.*, 1994 *J. Immunol.* 152, 4149, found that VEGF-specific antibodies were able to significantly reduce the mitogenic activity of synovial tissues from patients suffering from rheumatoid arthritis. These observations support a direct role for VEGF in rheumatoid 10 arthritis. Other angiogenic factors including those of the present invention may also be involved in arthritis.

5) Endometriosis: Various studies indicate that VEGF is directly implicated in endometriosis. In one study, VEGF concentrations measured by ELISA in peritoneal fluid were found to be significantly higher in women with endometriosis than in women without 15 endometriosis ( $24.1 \pm 15$  ng/ml vs  $13.3 \pm 7.2$  ng/ml in normals). In patients with endometriosis, higher concentrations of VEGF were detected in the proliferative phase of the menstrual cycle ( $33 \pm 13$  ng/ml) compared to the secretory phase ( $10.7 \pm 5$  ng/ml). The cyclic variation was not noted in fluid from normal patients (McLaren *et al.*, 1996, *Human Reprod.* 11, 220-223). In another study, women with moderate to severe endometriosis had significantly higher concentrations of peritoneal fluid VEGF than women without 20 endometriosis. There was a positive correlation between the severity of endometriosis and the concentration of VEGF in peritoneal fluid. In human endometrial biopsies, VEGF expression increased relative to the early proliferative phase approximately 1.6-, 2-, and 3.6-fold in midproliferative, late proliferative, and secretory endometrium (Shifren *et al.*, 1996, *J. Clin. Endocrinol. Metab.* 81, 3112-3118).

25 In a third study, VEGF-positive staining of human ectopic endometrium was shown to be localized to macrophages (double immunofluorescent staining with CD14 marker). Peritoneal fluid macrophages demonstrated VEGF staining in women with and without endometriosis. However, increased activation of macrophages (acid phosphatase activity) was demonstrated in fluid from women with endometriosis compared with controls. 30 Peritoneal fluid macrophage conditioned media from patients with endometriosis resulted in significantly increased cell proliferation ( $[^3\text{H}]$  thymidine incorporation) in HUVEC cells compared to controls. The percentage of peritoneal fluid macrophages with VEGFR2 mRNA was higher during the secretory phase, and significantly higher in fluid from women with endometriosis ( $80 \pm 15\%$ ) compared with controls ( $32 \pm 20\%$ ). Flt-mRNA was detected in

peritoneal fluid macrophages from women with and without endometriosis, but there was no difference between the groups or any evidence of cyclic dependence (McLaren *et al.*, 1996, *J. Clin. Invest.* 98, 482-489).

In the early proliferative phase of the menstrual cycle, VEGF has been found to be  
5 expressed in secretory columnar epithelium (estrogen-responsive) lining both the oviducts and the uterus in female mice. During the secretory phase, VEGF expression was shown to have shifted to the underlying stroma composing the functional endometrium. In addition to examining the endometrium, neovascularization of ovarian follicles and the corpus luteum, as well as angiogenesis in embryonic implantation sites have been analyzed. For these processes,  
10 VEGF was expressed in spatial and temporal proximity to forming vasculature (Shweiki *et al.*, 1993, *J. Clin. Invest.* 91, 2235-2243).

The present body of knowledge in VEGFR1 and/or VEGFR2 research indicates the need for methods to assay VEGFR1 and/or VEGFR2 activity and for compounds that can regulate VEGFR1 and/or VEGFR2 expression for research, diagnostic, and therapeutic use.  
15 As described herein, the nucleic acid molecules of the present invention can be used in assays to diagnose disease state related of VEGF, VEGFR1 and/or VEGFR2 levels. In addition, the nucleic acid molecules can be used to treat disease state related to VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 levels.

Particular processes, diseases, or conditions that can be associated with VEGFR1  
20 and/or VEGFR2 levels include, but are not limited to, gynecologic neovascularization, such as endometriosis, endometrial carcinoma, gynecologic bleeding disorders, irregular menstrual cycles, ovulation, premenstrual syndrome (PMS), menopausal dysfunction, other diseases and conditions discussed herein, and other diseases or conditions that are related to or respond to the levels of VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2, in a cell or tissue,  
25 alone or in combination with other therapies

The use of GnRH (gonadotropin releasing hormone) agonists, Lupron Depot (Leuprolide Acetate), Synarel (naferelin acetate), Zolodex (goserelin acetate), Suprefact (buserelin acetate), Danazol, or oral contraceptives including, but not limited to, Depo-Provera or Provera (medroxyprogesterone acetate), or any other estrogen/progesterone contraceptive, are  
30 all non-limiting examples of compounds and methods that can be combined with or used in conjunction with the nucleic acid molecules of the instant invention. Various chemotherapies can be readily combined with nucleic acid molecules of the invention for the treatment of endometrial carcinoma. Common chemotherapies that can be combined with nucleic acid molecules of the instant invention include various combinations of cytotoxic drugs to kill the

cancer cells. These drugs include but are not limited to paclitaxel (Taxol), docetaxel, cisplatin, methotrexate, cyclophosphamide, doxorubicin, fluorouracil carboplatin, edatrexate, gemcitabine, vinorelbine *etc.* Those skilled in the art will recognize that other drug compounds and therapies can be readily combined with the nucleic acid molecules of the 5 instant invention and are hence within the scope of the instant invention.

#### Animal Models

There are several animal models in which the anti-angiogenesis effect of nucleic acids of the present invention, such as ribozymes, directed against VEGF-R mRNAs can be tested. Typically, a corneal model has been used to study angiogenesis in rat and rabbit since 10 recruitment of vessels can easily be followed in this normally avascular tissue (Pandey *et al.*, 1995 *Science* 268: 567-569). In these models, a small Teflon or Hydron disk pretreated with an angiogenesis factor (e.g. bFGF or VEGF) is inserted into a pocket surgically created in the cornea. Angiogenesis is monitored 3 to 5 days later. Ribozymes directed against VEGF-R mRNAs would be delivered in the disk as well, or dropwise to the eye over the time course of 15 the experiment. In another eye model, hypoxia has been shown to cause both increased expression of VEGF and neovascularization in the retina (Pierce *et al.*, 1995 *Proc. Natl. Acad. Sci. USA.* 92: 905-909; Shweiki *et al.*, 1992 *J. Clin. Invest.* 91: 2235-2243).

In human glioblastomas, it has been shown that VEGF is at least partially responsible for tumor angiogenesis (Plate *et al.*, 1992 *Nature* 359, 845). Animal models have been 20 developed in which glioblastoma cells are implanted subcutaneously into nude mice and the progress of tumor growth and angiogenesis is studied (Kim *et al.*, 1993 *supra*; Millauer *et al.*, 1994 *supra*).

Another animal model that addresses neovascularization involves Matrigel, an extract of basement membrane that becomes a solid gel when injected subcutaneously (Passaniti *et al.*, 1992 *Lab. Invest.* 67: 519-528). When the Matrigel is supplemented with angiogenesis factors such as VEGF, vessels grow into the Matrigel over a period of 3 to 5 days and angiogenesis can be assessed. Ribozymes directed against VEGF-R mRNAs can be delivered 25 in the Matrigel to assess anti-angiogenesis effect.

Several animal models exist for screening of anti-angiogenic agents. These include 30 corneal vessel formation following corneal injury (Burger *et al.*, 1985 *Cornea* 4: 35-41; Lepri, *et al.*, 1994 *J. Ocular Pharmacol.* 10: 273-280; Ormerod *et al.*, 1990 *Am. J. Pathol.* 137: 1243-1252) or intracorneal growth factor implant (Grant *et al.*, 1993 *Diabetologia* 36: 282-291; Pandey *et al.* 1995 *supra*; Zieche *et al.*, 1992 *Lab. Invest.* 67: 711-715), vessel

growth into Matrigel matrix containing growth factors (Passaniti *et al.*, 1992 *supra*), female reproductive organ neovascularization following hormonal manipulation (Shweiki *et al.*, 1993 *Clin. Invest.* 91: 2235-2243), several models involving inhibition of tumor growth in highly vascularized solid tumors (O'Reilly *et al.*, 1994 *Cell* 79: 315-328; Senger *et al.*, 1993  
5 *Cancer and Metas. Rev.* 12: 303-324; Takahasi *et al.*, 1994 *Cancer Res.* 54: 4233-4237;  
Kim *et al.*, 1993 *supra*), and transient hypoxia-induced neovascularization in the mouse retina (Pierce *et al.*, 1995 *Proc. Natl. Acad. Sci. USA.* 92: 905-909).

The cornea model, described in Pandey *et al. supra*, is the most common and well characterized anti-angiogenic agent efficacy screening model. This model involves an avascular tissue into which vessels are recruited by a stimulating agent (growth factor, thermal or alkalai burn, endotoxin). The corneal model utilizes the intrastromal corneal implantation of a Teflon pellet soaked in a VEGF-Hydrone solution to recruit blood vessels toward the pellet which can be quantitated using standard microscopic and image analysis techniques. To evaluate their anti-angiogenic efficacy, ribozymes are applied topically to the eye or bound within Hydrone on the Teflon pellet itself. This avascular cornea as well as the Matrigel (see below) provide for low background assays. While the corneal model has been performed extensively in the rabbit, studies in the rat have also been conducted.  
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The mouse model (Passaniti *et al.*, *supra*) is a non-tissue model which utilizes Matrigel, an extract of basement membrane (Kleinman *et al.*, 1986) or Millipore® filter disk, which can be impregnated with growth factors and anti-angiogenic agents in a liquid form prior to injection. Upon subcutaneous administration at body temperature, the Matrigel or Millipore® filter disk forms a solid implant. VEGF embedded in the Matrigel or Millipore® filter disk would be used to recruit vessels within the matrix of the Matrigel or Millipore® filter disk which can be processed histologically for endothelial cell specific vWF (factor VIII antigen)  
20  
25  
30 immunohistochemistry, Trichrome-Masson stain, or hemoglobin content. Like the cornea, the Matrigel or Millipore® filter disk are avascular; however, it is not tissue. In the Matrigel or Millipore® filter disk model, ribozymes are administered within the matrix of the Matrigel or Millipore® filter disk to test their anti-angiogenic efficacy. Thus, delivery issues in this model, as with delivery of ribozymes by Hydrone-coated Teflon pellets in the rat cornea model, are minimized due to the homogeneous presence of the ribozyme within the respective matrix.

These models offer a distinct advantage over several other angiogenic models listed previously. The ability to use VEGF as a pro-angiogenic stimulus in both models is highly desirable since ribozymes target only VEGFr mRNA. In other words, the involvement of

other non-specific types of stimuli in the cornea and Matrigel models is not advantageous from the standpoint of understanding the pharmacologic mechanism by which the anti-VEGFr mRNA ribozymes produce their effects. In addition, the models allow for testing the specificity of the anti-VEGFr mRNA ribozymes by using either aFGF or bFGF as a pro-  
5 angiogenic factor. Vessel recruitment using FGF should not be affected in either model by anti-VEGFr mRNA ribozymes. Other models of angiogenesis, including vessel formation in the female reproductive system using hormonal manipulation (Shweiki *et al.*, 1993 *supra*); a variety of vascular solid tumor models which involve indirect correlations with angiogenesis  
10 (O'Reilly *et al.*, 1994 *supra*; Senger *et al.*, 1993 *supra*; Takahasi *et al.*, 1994 *supra*; Kim *et al.*, 1993 *supra*); and retinal neovascularization following transient hypoxia (Pierce *et al.*, 1995 *supra*), were not selected for efficacy screening due to their non-specific nature, although they can be useful models due to a demonstrated correlation between VEGF and angiogenesis.

Other model systems to study tumor angiogenesis is reviewed by Folkman, 1985 *Adv. 15 Cancer. Res.* 43, 175.

#### *Use of murine models*

For a typical systemic study involving 10 mice (20 g each) per dose group, 5 doses (1, 3, 10, 30 and 100 mg/kg daily over 14 days continuous administration), approximately 400 mg of ribozyme, formulated in saline would be used. A similar study in young adult rats (200 20 g) would require over 4 g. Parallel pharmacokinetic studies involve the use of similar quantities of ribozymes further justifying the use of murine models.

#### *Ribozymes and Lewis lung carcinoma and B-16 melanoma murine models*

Identifying a common animal model for systemic efficacy testing of ribozymes is an efficient way of screening ribozymes for systemic efficacy.

25 The Lewis lung carcinoma and B-16 murine melanoma models are well accepted models of primary and metastatic cancer and are used for initial screening of anti-cancer agents. These murine models are not dependent upon the use of immunodeficient mice, are relatively inexpensive, and minimize housing concerns. Both the Lewis lung and B-16 melanoma models involve subcutaneous implantation of approximately  $10^6$  tumor cells from 30 metastatically aggressive tumor cell lines (Lewis lung lines 3LL or D122, LLC-LN7; B-16-BL6 melanoma) in C57BL/6J mice. Alternatively, the Lewis lung model can be produced by the surgical implantation of tumor spheres (approximately 0.8 mm in diameter). Metastasis

also can be modeled by injecting the tumor cells directly intraveneously. In the Lewis lung model, microscopic metastases can be observed approximately 14 days following implantation with quantifiable macroscopic metastatic tumors developing within 21-25 days. The B-16 melanoma exhibits a similar time course with tumor neovascularization beginning 4 days following implantation. Since both primary and metastatic tumors exist in these models after 21-25 days in the same animal, multiple measurements can be taken as indices of efficacy. Primary tumor volume and growth latency as well as the number of micro- and macroscopic metastatic lung foci or number of animals exhibiting metastases can be quantitated. The percent increase in lifespan can also be measured. Thus, these models provide suitable primary efficacy assays for screening systemically administered ribozymes/ribozyme formulations.

In the Lewis lung and B-16 melanoma models, systemic pharmacotherapy with a wide variety of agents usually begins 1-7 days following tumor implantation/inoculation with either continuous or multiple administration regimens. Concurrent pharmacokinetic studies can be performed to determine whether sufficient tissue levels of ribozymes can be achieved for pharmacodynamic effect to be expected. Furthermore, primary tumors and secondary lung metastases can be removed and subjected to a variety of *in vitro* studies (*i.e.* target RNA reduction).

Flt-1, KDR and/or flk-1 protein levels can be measured clinically or experimentally by FACS analysis. Flt-1, KDR and/or flk-1 encoded mRNA levels can be assessed by Northern analysis, RNase-protection, primer extension analysis and/or quantitative RT-PCR. Ribozymes that block flt-1, KDR and/or flk-1 protein encoding mRNAs and therefore result in decreased levels of flt-1, KDR and/or flk-1 activity by more than 20% *in vitro* can be identified.

Ribozymes and/or genes encoding them are delivered by either free delivery, liposome delivery, cationic lipid delivery, adeno-associated virus vector delivery, adenovirus vector delivery, retrovirus vector delivery or plasmid vector delivery in these animal model experiments (see above).

Subjects can be treated by locally administering nucleic acids targeted against VEGF-R by direct injection. Routes of administration include, but are not limited to, intravascular, intramuscular, subcutaneous, intraarticular, aerosol inhalation, oral (tablet, capsule or pill form), topical, systemic, ocular, intraperitoneal and/or intrathecal delivery.

Surgically induced models of endometriosis have been developed in rats, mice, and rabbits. Non-human primates demonstrate spontaneous endometriosis, but surgical induction can also be used. In addition to the surgical technique, cycle monitoring can be performed by daily vaginal cytology in primates. For all of the surgically induced models of endometriosis,

5 the following general procedure is used. An initial laparotomy is performed to implant tissue from a donor animal. A portion of one uterine horn (or one complete horn in the case of mice) is removed. The endometrium of this piece of uterus is separated from the myometrium and cut into small segments (4-10 mm<sup>2</sup>). Segments (approximately 3) are sutured to various locations within the abdominal cavity (peritoneum, intestinal mesentery vessels, uterus, broad

10 ligament). Cummings and Metcalf (1996) attached whole segments of mouse uterus without separating the endometrium from the myometrium. Implants are allowed to grow for 3-6 weeks. A second laparotomy is sometimes performed to verify development of endometriosis-like foci (vascularization and cysts filled with clear fluid). This second laparotomy was done in the studies by Quereda *et al.*, (1996) and Stoeckemann *et al.*, (1995).

15 After 3-6 weeks post-surgery and/or following visualization of endometriosis, drug treatment is initiated and continued for a prescribed period of time. At the termination of these studies, animals are euthanized. Endpoints include, but are not limited to, changes in the surface area of the implants and tissue mass of the ectopic endometrial implants (see for example Brogniez *et al.*, 1995, *Human Reprod.* 10, 927-931; Cummings *et al.*, 1996, *Tox. Appl. Pharm.* 138, 131-139; Cummings and Metcalf, 1996, *Proc. Soc. Exp. Biol. Med.* 212, 332-

20 337; D'Hooghe *et al.*, 1996, *Fertility and Sterility.* 66, 809-813; Quereda *et al.*, 1996, *Eur. J. Obstet. Gynecol. Rep. Biol.* 67, 35-40; and Stoeckemann *et al.*, 1995, *Human Reprod.* 10, 3264-3271).

#### *Combination therapies*

25 Gemcytabine and cyclophosphamide are non-limiting examples of chemotherapeutic agents that can be combined with or used in conjunction with the nucleic acid molecules (e.g. ribozymes and antisense molecules) of the instant invention. Those skilled in the art will recognize that other anti-angiogenic and/or anti-cancer compounds and therapies can be similarly be readily combined with the nucleic acid molecules of the instant invention (e.g. 30 ribozymes and antisense molecules) and are hence within the scope of the instant invention. Such compounds and therapies are well known in the art (see for example Cancer: Principles

and Practice of Oncology, Volumes 1 and 2, eds Devita, V.T., Hellman, S., and Rosenberg, S.A., J.B. Lippincott Company, Philadelphia, USA; incorporated herein by reference) and include, without limitations, folates, antifolates, pyrimidine analogs, fluoropyrimidines, purine analogs, adenosine analogs, topoisomerase I inhibitors, anthrapyrazoles, retinoids, antibiotics, anthacyclins, platinum analogs, alkylating agents, nitrosoureas, plant derived compounds such as vinca alkaloids, epipodophyllotoxins, tyrosine kinase inhibitors, taxols, radiation therapy, surgery, nutritional supplements, gene therapy, radiotherapy, for example 3D-CRT, immunotoxin therapy, for example ricin, and monoclonal antibodies. Specific examples of chemotherapeutic compounds than can be combined with or used in conjunction with the nucleic acid molecules of the invention include but are not limited to Paclitaxel; Docetaxel; Methotrexate; Doxorubicin; Edatrexate; Vinorelbine; Tomoxifen; Leucovorin; 5-fluoro uridine (5-FU); Irinotecan (CAMPTOSAR® or CPT-11 or Camptothecin-11 or Campto); Cisplatin; Carboplatin; Amsacrine; Cytarabine; Bleomycin; Mitomycin C; Dactinomycin; Mithramycin; Hexamethylmelamine; Dacarbazine; L-asperginate; Nitrogen mustard; Melphalan, Chlorambucil; Busulfan; Ifosfamide; 4-hydroperoxycyclophosphamide, Thiotepa; Tamoxifen, Herceptin; IMC C225; ABX-EGF; and combinations thereof.

#### Diagnostic uses

The nucleic acid molecules of this invention (e.g., enzymatic nucleic acid molecules) can be used as diagnostic tools to examine genetic drift and mutations within diseased cells or to detect the presence of VEGF and/or VEGFr, such as VEGFR1 and/or VEGFR2 RNA in a cell. The close relationship between enzymatic nucleic acid molecule activity and the structure of the target RNA allows the detection of mutations in any region of the molecule which alters the base-pairing and three-dimensional structure of the target RNA. By using multiple enzymatic nucleic acid molecules described in this invention, one can map nucleotide changes which are important to RNA structure and function *in vitro*, as well as in cells and tissues. Cleavage of target RNAs with enzymatic nucleic acid molecules can be used to inhibit gene expression and define the role (essentially) of specified gene products in the progression of disease. In this manner, other genetic targets can be defined as important mediators of the disease. These experiments can lead to better treatment of the disease progression by affording the possibility of combinational therapies (e.g., multiple enzymatic nucleic acid molecules targeted to different genes, enzymatic nucleic acid molecules coupled with known small molecule inhibitors, or intermittent treatment with combinations of enzymatic nucleic acid molecules and/or other chemical or biological molecules). Other *in*

*vitro* uses of enzymatic nucleic acid molecules of this invention are well known in the art, and include detection of the presence of mRNAs associated with VEGF, VEGFR1 and/or VEGFR2-related condition. Such RNA is detected by determining the presence of a cleavage product after treatment with an enzymatic nucleic acid molecule using standard methodology.

5        In a specific example, enzymatic nucleic acid molecules which cleave only wild-type or mutant forms of the target RNA are used for the assay. The first enzymatic nucleic acid molecule is used to identify wild-type RNA present in the sample and the second enzymatic nucleic acid molecule is used to identify mutant RNA in the sample. As reaction controls, synthetic substrates of both wild-type and mutant RNA are cleaved by both enzymatic nucleic  
10      acid molecules to demonstrate the relative enzymatic nucleic acid molecule efficiencies in the reactions and the absence of cleavage of the "non-targeted" RNA species. The cleavage products from the synthetic substrates also serve to generate size markers for the analysis of wild-type and mutant RNAs in the sample population. Thus each analysis requires two enzymatic nucleic acid molecules, two substrates and one unknown sample which is  
15      combined into six reactions. The presence of cleavage products is determined using an RNase protection assay so that full-length and cleavage fragments of each RNA can be analyzed in one lane of a polyacrylamide gel. It is not absolutely required to quantify the results to gain insight into the expression of mutant RNAs and putative risk of the desired phenotypic changes in target cells. The expression of mRNA whose protein product is  
20      implicated in the development of the phenotype (*i.e.*, VEGFR1 and/or VEGFR2) is adequate to establish risk. If probes of comparable specific activity are used for both transcripts, then a qualitative comparison of RNA levels will be adequate and will decrease the cost of the initial diagnosis. Higher mutant form to wild-type ratios are correlated with higher risk whether RNA levels are compared qualitatively or quantitatively. The use of enzymatic nucleic acid  
25      molecules in diagnostic applications contemplated by the instant invention is described, for example, in Usman *et al.*, US Patent Application No. 09/877,526, George *et al.*, US Patent Nos. 5,834,186 and 5,741,679, Shih *et al.*, US Patent No. 5,589,332, Nathan *et al.*, US Patent No 5,871,914, Nathan and Ellington, International PCT publication No. WO 00/24931, Breaker *et al.*, International PCT Publication Nos. WO 00/26226 and 98/27104, and  
30      Sullenger *et al.*, US Patent Application Serial No. 09/205,520.

#### Additional Uses

Uses of sequence-specific enzymatic nucleic acid molecules of the instant invention can have many of the same applications for the study of RNA that DNA restriction endonucleases have for the study of DNA (Nathans *et al.*, 1975 *Ann. Rev. Biochem.* 44:273). For example,

the pattern of restriction fragments can be used to establish sequence relationships between two related RNAs, and large RNAs can be specifically cleaved to fragments of a size more useful for study. The ability to engineer sequence specificity of the enzymatic nucleic acid molecule is ideal for cleavage of RNAs of unknown sequence. Applicant has described the  
5 use of nucleic acid molecules to down-regulate gene expression of target genes in bacterial, microbial, fungal, viral, and eukaryotic systems including plant, or mammalian cells.

All patents and publications mentioned in the specification are indicative of the levels of skill of those skilled in the art to which the invention pertains. All references cited in this disclosure are incorporated by reference to the same extent as if each reference had been  
10 incorporated by reference in its entirety individually.

One skilled in the art would readily appreciate that the present invention is well adapted to carry out the objects and obtain the ends and advantages mentioned, as well as those inherent therein. The methods and compositions described herein as presently representative of preferred embodiments are exemplary and are not intended as limitations on the scope of  
15 the invention. Changes therein and other uses will occur to those skilled in the art, which are encompassed within the spirit of the invention, are defined by the scope of the claims.

It will be readily apparent to one skilled in the art that varying substitutions and modifications may be made to the invention disclosed herein without departing from the scope and spirit of the invention. Thus, such additional embodiments are within the scope of  
20 the present invention and the following claims.

The invention illustratively described herein suitably may be practiced in the absence of any element or elements, limitation or limitations which is not specifically disclosed herein. Thus, for example, in each instance herein any of the terms "comprising", "consisting essentially of" and "consisting of" may be replaced with either of the other two terms. The  
25 terms and expressions which have been employed are used as terms of description and not of limitation, and there is no intention that in the use of such terms and expressions of excluding any equivalents of the features shown and described or portions thereof, but it is recognized that various modifications are possible within the scope of the invention claimed. Thus, it should be understood that although the present invention has been specifically disclosed by  
30 preferred embodiments, optional features, modification and variation of the concepts herein disclosed may be resorted to by those skilled in the art, and that such modifications and variations are considered to be within the scope of this invention as defined by the description and the appended claims.

In addition, where features or aspects of the invention are described in terms of Markush groups or other grouping of alternatives, those skilled in the art will recognize that the invention is also thereby described in terms of any individual member or subgroup of members of the Markush group or other group.

- 5        Other embodiments are within the following claims.

TABLE ICharacteristics of Ribozymes**Group I Introns**

Size: ~200 to >1000 nucleotides.

Requires a U in the target sequence immediately 5' of the cleavage site.

Binds 4-6 nucleotides at 5' side of cleavage site.

Over 75 known members of this class. Found in *Tetrahymena thermophila* rRNA, fungal mitochondria, chloroplasts, phage T4, blue-green algae, and others.

**RNaseP RNA (M1 RNA)**

Size: ~290 to 400 nucleotides.

RNA portion of a ribonucleoprotein enzyme. Cleaves tRNA precursors to form mature tRNA.

Roughly 10 known members of this group all are bacterial in origin.

**Hammerhead Ribozyme**

Size: ~13 to 40 nucleotides.

Requires the target sequence UH immediately 5' of the cleavage site.

Binds a variable number of nucleotides on both sides of the cleavage site.

14 known members of this class. Found in a number of plant pathogens (virusoids) that use RNA as the infectious agent (Figure 1 and 2)

**Hairpin Ribozyme**

Size: ~50 nucleotides.

Requires the target sequence GUC immediately 3' of the cleavage site.

Binds 4-6 nucleotides at 5' side of the cleavage site and a variable number to the 3' side of the cleavage site.

Only 3 known member of this class. Found in three plant pathogen (satellite RNAs of the tobacco ringspot virus, arabis mosaic virus and chicory yellow mottle virus) which uses RNA as the infectious agent (Figure 3).

**Hepatitis Delta Virus (HDV) Ribozyme**

Size: 50 - 60 nucleotides (at present).

Sequence requirements not fully determined.

Binding sites and structural requirements not fully determined, although no sequences 5' of cleavage site are required.

Only 1 known member of this class. Found in human HDV (Figure 4).

***Neurospora* VS RNA Ribozyme**

**Size: ~144 nucleotides (at present)**

**Cleavage of target RNAs recently demonstrated.**

**Sequence requirements not fully determined.**

**Binding sites and structural requirements not fully determined. Only 1 known member of this class. Found in *Neurospora* VS RNA (Figure 5).**

**Table II:****A. 2.5 μmol Synthesis Cycle ABI 394 Instrument**

<b>Reagent</b>	<b>Equivalents</b>	<b>Amount</b>	<b>Wait Time*</b>	<b>DNA</b>	<b>Wait Time* 2'-O-methyl</b>	<b>Wait Time*</b>
<b>Phosphoramidites</b>	6.5	163 μL	45 sec	2.5 min	7.5 min	
S-Ethyl Tetrazole	23.8	238 μL	45 sec	2.5 min	7.5 min	
Acetic Anhydride	100	233 μL	5 sec	5 sec	5 sec	
N-Methyl Imidazole	186	233 μL	5 sec	5 sec	5 sec	
TCA	176	2.3 mL	21 sec	21 sec	21 sec	
Iodine	11.2	1.7 mL	45 sec	45 sec	45 sec	
Beaucage	12.9	645 μL	100 sec	300 sec	300 sec	
Acetonitrile	NA	6.67 mL	NA	NA	NA	

**B. 0.2 μmol Synthesis Cycle ABI 394 Instrument**

<b>Reagent</b>	<b>Equivalents</b>	<b>Amount</b>	<b>Wait Time*</b>	<b>DNA</b>	<b>Wait Time* 2'-O-methyl</b>	<b>Wait Time*</b>
<b>Phosphoramidites</b>	15	31 μL	45 sec	233 sec	465 sec	
S-Ethyl Tetrazole	38.7	31 μL	45 sec	233 min	465 sec	
Acetic Anhydride	655	124 μL	5 sec	5 sec	5 sec	
N-Methyl Imidazole	1245	124 μL	5 sec	5 sec	5 sec	
TCA	700	732 μL	10 sec	10 sec	10 sec	
Iodine	20.6	244 μL	15 sec	15 sec	15 sec	

Beaucage	7.7	232 $\mu$ L	100 sec	300 sec	300 sec
Acetonitrile	NA	2.64 mL	NA	NA	NA

**C. 0.2  $\mu$ mol Synthesis Cycle 96 well Instrument**

Reagent	Equivalents DNA/2'-O-methyl/Ribonucleotide	Amount DNA/2'-O-methyl/Ribonucleotide	Wait Time* DNA	Wait Time* 2'-O-methyl methyl	Wait Time* Ribonucleotide
Phosphoramidites	22/33/66	40/60/120 $\mu$ L	60 sec	180 sec	360 sec
S-Ethyli Tetrazole	70/105/210	40/60/120 $\mu$ L	60 sec	180 min	360 sec
Acetic Anhydride	265/265/265	50/50/50 $\mu$ L	10 sec	10 sec	10 sec
N-Methyl Imidazole	502/502/502	50/50/50 $\mu$ L	10 sec	10 sec	10 sec
TCA	236/475/475	250/500/500 $\mu$ L	15 sec	15 sec	15 sec
Iodine	6.8/6.8/6.8	80/80/80 $\mu$ L	30 sec	30 sec	30 sec
Beaucage	34/51/51	80/120/120	100 sec	200 sec	200 sec
Acetonitrile	NA	1150/1150/1150 $\mu$ L	NA	NA	NA

\* Wait time does not include contact time during delivery.

**Table III: Patient Demographics**

Dose cohort (mg/m <sup>2</sup> )	Pt#	Age	Sex	Diagnosis	Doses
10	1001	49	F	NSC Lung	29
10	1002	65	F	liposarcoma	120
10	1003	49	M	nasopharyngeal CA	109
30	1004	35	M	non-small cell lung	1
30	1005	45	F	melanoma (ocular)	113
30	1006	57	M	colon	199
30	1007	39	F	epithelioid hemangioendothelioma	198
100	1008	52	M	adrenal CA	57
100	1009	44	F	breast	35
100	1010	62	F	renal	134
300	1011	24	F	melanoma	31
300	1012	57	M	renal cell	178
300	1013	53	M	nasopharyngeal SCCA	29
300	1014	64	F	peritoneal mesothelioma	324
100	1015	65	M	melanoma	140
100	1016	77	F	breast	265
100	1017		F	melanoma	35
100	1018	26	F	melanoma	7
100	1019	69	F	endometrial sarcoma	500
100	1020	65	M	carcinoid	124
100	1021	59	M	gallbladder adeno carcinoma	34
100	1022	43	M	colorectal	8
100	1023	78	F	breast	50
100	1024	40	F	parotid adenocarcinoma	285
100	1025	52	F	breast	71
100	1026	39	F	breast	34
100	1027	55	F	breast	36
100	1028	52	M	melanoma	29
100	1029	38	M	pancreatic	36
100	1030	83	M	melanoma	41
100	1031	50	M	medullary thyroid	108

One patient taken off study due to progressive disease. Allowed to resume ANGIOZYME on a compassionate basis.

As of September 1, 2001, all patients were off study. (Although one patient resumed treatment per above note)

Table IV Pharmacokinetic parameters of ANGIOZYME after bolus subcutaneous administration.

	10 mg/m <sup>2</sup>		30 mg/m <sup>2</sup>		100 mg/m <sup>2</sup>		300 mg/m <sup>2</sup>	
	Mean	SD	Mean	SD	Mean	SD	Mean	SD
<b>Day 1</b>								
Cmax (ug/mL)	0.43	0.07	0.62	0.28	3.17	0.69	8.91	2.93
AUCt (ug*hr/mL)	2.60	1.43	6.04	2.70	34.14	2.28	89.87	21.68
AUCinf (ug*hr/mL)	4.40	0.06	7.99	1.66	37.51	1.91	101.57	13.47
t(1/2) (hr)	3.62	0.79	7.32	6.94	4.58	0.02	9.26	6.20
CL/F (L/hr/m <sup>2</sup> )	2.24	0.08	3.73	0.92	2.96	0.61	2.99	0.43
<b>Day 29</b>								
Cmax (ug/mL)	0.35	0.19	1.17	0.53	3.23	0.35	8.93	6.71
AUCt (ug*hr/mL)	2.11	1.31	7.29	1.16	31.87	1.91	119.42	65.84
AUCinf (ug*hr/mL)	3.38	1.31	8.54	2.46	33.61	2.16	132.73	67.82
t(1/2) (hr)	4.49	1.60	3.26	1.01	4.66	0.35	7.24	0.70
CL/F (L/hr/m <sup>2</sup> )	2.49	1.48	3.69	0.94	3.21	0.56	2.72	1.40

Table V: Human FLT DNAzyme and Substrate Sequence

Pos	Substrate	Seq ID No	DNAzyme	Seq ID No
17	UCCUCUCG G CUCCUCCC	1	GGGAGGAG GGCTAGCTACAACGA CGAGAGGA	1703
28	CCUCCCCG G CAGCGGCG	2	CGCCGCTG GGCTAGCTACAACGA CGGGGAGG	1704
31	CCCCGGCA G CGGGGGCG	3	CGCCGCCG GGCTAGCTACAACGA TGCCGGGG	1705
34	CGGCAGCG G CGGCGGCU	4	AGCCGCCG GGCTAGCTACAACGA CGCTGCCG	1706
37	CAGCGGGC G CGGCUCGG	5	CCGAGCCG GGCTAGCTACAACGA CGCCGCTG	1707
40	CGGCGGGC G CUoggGAC	6	GCTCCGAG GGCTAGCTACAACGA CGCCGCCG	1708
47	GGCUCGGA G CGGGCUC	7	GGAGCCCG GGCTAGCTACAACGA TCCGAGCC	1709
51	CGGAGCGG G CUCCGGGG	8	CCCCGGAG GGCTAGCTACAACGA CCGCTCCG	1710
59	GCUCGGGG G CUCGGGUG	9	CACCCGAG GGCTAGCTACAACGA CCCGGAGC	1711
65	GGGCUCGG G UGGCAGGG	10	CCGCTGCA GGCTAGCTACAACGA CCGAGCCC	1712
67	GCUCGGGU G CAGCGGCC	11	GGCCGCTG GGCTAGCTACAACGA ACCCGAGC	1713
70	CGGGUGCA G CGGCCAGC	12	GCTGGCCG GGCTAGCTACAACGA TGCACCCG	1714
73	GUGCACCG G CCAGCGGG	13	CCCGCTGG GGCTAGCTACAACGA CGCTGAC	1715
77	AGCGGGCCA G CGGGCCUG	14	CAGGCCCG GGCTAGCTACAACGA TGCCGGCT	1716
81	GCCAGCGG G CCUGGCGG	15	CCGCCAGG GGCTAGCTACAACGA CCGCTGGC	1717
86	CGGGCCUG G CGGCGAGG	16	CCTCGCCG GGCTAGCTACAACGA CAGGCCCG	1718
89	GCUUGGCG G CGAGGAUU	17	AATCCTCG GGCTAGCTACAACGA CGCCAGGC	1719
95	CGGGCGAG A UUACCCGG	18	CCGGGTAA GGCTAGCTACAACGA CCTCGCCG	1720
98	CGAGGAUT A CCCGGGGA	19	TCCCGGGG GGCTAGCTACAACGA AATCCTCG	1721
108	CGGGGGAA G UGGUUGUC	20	GACAACCA GGCTAGCTACAACGA TTCCCCGG	1722
111	GGGAAGUG G UUGUCUCC	21	GGAGACAA GGCTAGCTACAACGA CACTTCCC	1723
114	AAGUGGUU G UCUCUCCUG	22	CCAGGAGA GGCTAGCTACAACGA AACCACTT	1724
122	GUUCUCUG G CUGGAGCC	23	GGCTCCAG GGCTAGCTACAACGA CAGGAGAC	1725
128	UGGGUGGA G CCGCGAGA	24	TCTCGCCG GGCTAGCTACAACGA TCCAGCCA	1726
131	CUGGAGCC G CGAGACGG	25	CCGTCTCG GGCTAGCTACAACGA GGCTCCAG	1727
136	GCCGCGAG A CGGGCGCU	26	AGCGCCCG GGCTAGCTACAACGA CTCCGCCG	1728
140	CGAGACGG G CGCUCAGG	27	CCTGAGCG GGCTAGCTACAACGA CCGTCTCG	1729
142	AGACGGGC G CUCAGGGC	28	GCCCTGAG GGCTAGCTACAACGA GCCCGTCT	1730
149	CGCUCAGG G CGGGGGGC	29	GCCCCGCG GGCTAGCTACAACGA CCTGAGCG	1731
151	CUCAGGGC G CGGGCGCG	30	CGGCCCGG GGCTAGCTACAACGA GCCCTGAG	1732
156	GCGCGGGG G CGGGGGGC	31	GCCGCCGG GGCTAGCTACAACGA CCCGCGCC	1733
160	CGGGGGCG G CGGGGGCG	32	CGGCCCGG GGCTAGCTACAACGA CGGGCCCG	1734
163	GCGCGGGG G CGGCCAAC	33	GTTGCCCC GGCTAGCTACAACGA CGGGGGCC	1735
166	CGGGGGCG G CGAACCGAG	34	CTCGTTCG GGCTAGCTACAACGA CGGGGCCG	1736
170	GGCGGGCG A CGAGAGGA	35	TCCCTCTCG GGCTAGCTACAACGA TCGCCGCC	1737
178	ACGAGAGG A CGGACUCU	36	AGAGTCGG GGCTAGCTACAACGA CCTCTCGT	1738
182	GAGGACGG A CUCUGGCG	37	CGCCAGAG GGCTAGCTACAACGA CGTCCTC	1739
188	GGACUCUG G CGGGGGGG	38	CCCGGCCG GGCTAGCTACAACGA CAGAGTCC	1740
191	CUCUGGCG G CGGGGGCG	39	CGACCCGG GGCTAGCTACAACGA CGCCAGAG	1741
196	GCGGCCGG G UCGUUGGC	40	GCCAACGA GGCTAGCTACAACGA CGGGCCGC	1742
199	GCGGGGUC G UGGGGCGG	41	CCGGCCAA GGCTAGCTACAACGA GACCCGGC	1743
203	GGUCGUUG G CGGGGGGA	42	TCCCCCGG GGCTAGCTACAACGA CAACGACC	1744
212	CGGGGGGA G CGGGGGCA	43	TGCCCCGG GGCTAGCTACAACGA TCCCCGGG	1745
214	GGGGGAGC G CGGGGCAAC	44	GGTGCCCC GGCTAGCTACAACGA GCTCCCCC	1746
218	GAGCGCGG G CACCGGGC	45	GCCCGGTG GGCTAGCTACAACGA CGCGCTC	1747
220	GCGCGGGC A CGGGGGCA	46	TCGCCCCG GGCTAGCTACAACGA GCGCGC	1748
225	GGCACCGG G CGAGCAGG	47	CCTGCTCG GGCTAGCTACAACGA CGGGTGCC	1749
229	CGGGGGCGA G CAGGCCGC	48	GCGGCCCTG GGCTAGCTACAACGA TCGCCCGG	1750

233	GCGAGCAG G CGCGCGUCG	49	CGACGCGG GGCTAGCTACAACGA CTGCTCGC	1751
236	AGCAGGCC G CGUCGCGC	50	GCGCGACG GGCTAGCTACAACGA GGCCCTGCT	1752
238	CAGGCCGC G UCGCGCUC	51	GAGCGCGA GGCTAGCTACAACGA GCGGCCTG	1753
241	GCGCGCUC G CGCUCACC	52	GGTGAGCG GGCTAGCTACAACGA GACGCCGC	1754
243	CGCGUCGC G CUCACCAU	53	ATGGTGAG GGCTAGCTACAACGA GCGACGCG	1755
247	UCGCGCUC A CCAUGGUC	54	GACCATGG GGCTAGCTACAACGA GAGGCGGA	1756
250	CGCUCACC A UGGUCAGC	55	GCTGACCA GGCTAGCTACAACGA GGTGACCG	1757
253	UCACCAUG G UCAGCUAC	56	GTAGCTGA GGCTAGCTACAACGA CATGGTGA	1758
257	CAUGGUCA G CUACUGGG	57	CCCAGTAG GGCTAGCTACAACGA TGACCATG	1759
260	GGUCAGCU A CUUUGGACA	58	TGTCCCCAG GGCTAGCTACAACGA AGCTGACC	1760
266	CUACUGGG A CACCGGGG	59	CCCCGGTG GGCTAGCTACAACGA CCCAGTAG	1761
268	ACUGGGAC A CGGGGGUC	60	GACCCCGG GGCTAGCTACAACGA GTCCCCAGT	1762
274	ACACCGGG G UCCUGCUG	61	CAGCAGGA GGCTAGCTACAACGA CCCGGTGT	1763
279	GGGGGUCCU G CUGUGCGC	62	GCGCACAG GGCTAGCTACAACGA AGGACCCC	1764
282	GUCCUGCU G UGGCGCGU	63	AGCGCGCA GGCTAGCTACAACGA AGCAGGAC	1765
284	CCUGCUGU G CGCGCUGC	64	GCAGCGCG GGCTAGCTACAACGA ACAGCAGG	1766
286	UGCUGUGC G CGCUGCUC	65	GAGCAGCG GGCTAGCTACAACGA GCACAGCA	1767
288	CUGUGOGC G CUGCUCAG	66	CTGAGCAG GGCTAGCTACAACGA GCGCACAG	1768
291	UGCGCGCU G CUCAGCUG	67	CAGCTGAG GGCTAGCTACAACGA AGCGCGCA	1769
296	GCUGCUCA G CUGUCUGC	68	GCAGACAG GGCTAGCTACAACGA TGAGCAGC	1770
299	GCUCAGCU G UCUGCUUC	69	GAAGCAGA GGCTAGCTACAACGA AGCTGACC	1771
303	AGCUGUCU G CUUCUCAC	70	GTGAGAAG GGCTAGCTACAACGA AGACAGCT	1772
310	UGCUUCUC A CAGGAUCU	71	AGATCTG GGCTAGCTACAACGA GAGAACCA	1773
315	CUCACAGG A UCUAGUUC	72	GAACTAGA GGCTAGCTACAACGA CCTGTGAG	1774
320	AGGAUCUA G UUCAGGUU	73	AACCTGAA GGCTAGCTACAACGA TAGATCCT	1775
326	UAGUUCAG G UUCAAAA	74	ATTTTGAA GGCTAGCTACAACGA CTGAACTA	1776
333	GGUUCAAA A UUAAAAGA	75	TCTTTTAA GGCTAGCTACAACGA TTTGAACC	1777
341	AUAAAAG A UCCUGAAC	76	GTTCAGGA GGCTAGCTACAACGA CTTTTAAT	1778
348	GAUCCUGA A CUGAGUUU	77	AAACTCAG GGCTAGCTACAACGA TCAGGGTC	1779
353	UGAACUGA G UUUTAAAAG	78	CTTTTAAA GGCTAGCTACAACGA TCAGTTCA	1780
362	UUUTAAAAG G CACCCAGC	79	GCTGGGTG GGCTAGCTACAACGA CTTTTAAA	1781
364	UAAAAGGC A CCCAGCAC	80	GTGCTGGG GGCTAGCTACAACGA GCCTTTTA	1782
369	GGCACCCA G CACAUCAU	81	ATGATGTG GGCTAGCTACAACGA TGGGTGCC	1783
371	CACCCAGC A CAUCAUGC	82	GCATGATG GGCTAGCTACAACGA GCTGGGTG	1784
373	CCCAGCAC A UCAUGCAA	83	TTGCATGA GGCTAGCTACAACGA GTGCTGGG	1785
376	AGCACAUC A UGCAAGCA	84	TGCTTGCA GGCTAGCTACAACGA GATGTGCT	1786
378	CACAUCAU G CAAGCAGG	85	CCTGTTG GGCTAGCTACAACGA ATGATGTG	1787
382	UCAUGCAA G CAGGCCAG	86	CTGGCTG GGCTAGCTACAACGA TTGCATGA	1788
386	GCAAGCAG G CCAGACAC	87	GTGTCCTGG GGCTAGCTACAACGA CTGCTTGC	1789
391	CAGGCCAG A CACUGCAU	88	ATGCAGTG GGCTAGCTACAACGA CTGGCTG	1790
393	GGCCAGAC A CUGCAUTU	89	AGATGCCAG GGCTAGCTACAACGA GTCTGGCC	1791
396	CAGACACU G CAUCUCCA	90	TGGAGATG GGCTAGCTACAACGA AGTGTCTG	1792
398	GACACUGC A UCUCCAAU	91	ATTGGAGA GGCTAGCTACAACGA GCAGTGTC	1793
405	CAUCUCCA A UGCAGGGG	92	CCCCTGCA GGCTAGCTACAACGA TGGAGATG	1794
407	UCUCCAAU G CAGGGGG	93	CCCCCTG GGCTAGCTACAACGA ATTGGAGA	1795
418	GGGGGGAA G CAGCCCAU	94	ATGGGCTG GGCTAGCTACAACGA TTCCCCCC	1796
421	GGGAAGCA G CCCAUAAA	95	TTTATGGG GGCTAGCTACAACGA TGCTTCCC	1797
425	AGCAGCCC A UAAAUGGU	96	ACCATTGA GGCTAGCTACAACGA GGGCTGCT	1798
429	GCCCAUAA A UGGUCUUU	97	AAAGACCA GGCTAGCTACAACGA TTATGGC	1799
432	CAUAAAUG G UCUUUGCC	98	GGCAAAGA GGCTAGCTACAACGA CATTTATG	1800
438	UGGUCUUU G CCUGAAAU	99	ATTCAGG GGCTAGCTACAACGA AAAGACCA	1801
445	UGCCUGAA A UGGUGAGU	100	ACTCACCA GGCTAGCTACAACGA TTCAGGCA	1802

448	CUGAAAUG G UGAGUAAAG	101	CTTACTCA GGCTAGCTACAACGA CATTTCAG	1803
452	AAUGGUGA G UAAGGAAA	102	TTTCCCTTA GGCTAGCTACAACGA TCACCATT	1804
461	UAAGGAAA G CGAAAGGC	103	GCCTTTCG GGCTAGCTACAACGA TTTCCTTA	1805
468	AGCGAAAG G CUGAGCAU	104	ATGCTCAG GGCTAGCTACAACGA CTTTCGCT	1806
473	AAGGCUGA G CAUACUCA	105	TAGTTATG GGCTAGCTACAACGA TCAGCCCT	1807
475	GGCUGAGC A UAACUAAA	106	TTTAGTTA GGCTAGCTACAACGA GCTCAGCC	1808
478	UGAGCAUA A CUAAAUCU	107	AGATTTAG GGCTAGCTACAACGA TATGCTCA	1809
483	AUAACUAA A UCUGCCUG	108	CAGGCAGA GGCTAGCTACAACGA TTAGTTAT	1810
487	CUAAAUCU G CCUGUGGA	109	TCCACAGG GGCTAGCTACAACGA AGATTTAG	1811
491	AUCUGCCU G UGGAAGAA	110	TTCTTCCA GGCTAGCTACAACGA AGGCAGAT	1812
500	UGGAAGAA A UGGCAAAAC	111	GTTTGCCG GGCTAGCTACAACGA TTCTTCCA	1813
503	AAGAAAUG G CAAACAAU	112	ATTGTTTG GGCTAGCTACAACGA CATTCTT	1814
507	AAUGGCAA A CAUUCUG	113	CAGAATTG GGCTAGCTACAACGA TTGCCATT	1815
510	GGCAAACAA C UUCUGCAG	114	CTGCAGAA GGCTAGCTACAACGA TGTTTGCC	1816
515	ACAAUUCU G CAGUACUU	115	AAGTACTG GGCTAGCTACAACGA AGAATTGT	1817
518	AUUCUGCA G UACUUUAA	116	TTAAAGTA GGCTAGCTACAACGA TGCAGAAT	1818
520	UCUGCAGU A CUUUAACC	117	GGTTAAAG GGCTAGCTACAACGA ACTGCAGA	1819
526	GUACUUUA A CCUUGAAC	118	GTTCAAGG GGCTAGCTACAACGA TAAAGTAC	1820
533	AACCUUGA A CACAGCUC	119	GAGCTGTG GGCTAGCTACAACGA TCAAGGTT	1821
535	CCUUGAAC A CAGCUCAA	120	TTGAGCTG GGCTAGCTACAACGA GTTCAAGG	1822
538	UGAACACAA G CUCUAGCA	121	TGCTTGAG GGCTAGCTACAACGA TGTGTTCA	1823
544	CAGCUCAA G CAAACCCAC	122	GTGGTTTG GGCTAGCTACAACGA TTGAGCTG	1824
548	UCAAGCAA A CCACACUG	123	CAGTGTGG GGCTAGCTACAACGA TTGCTTGA	1825
551	AGCAAACC A CACUGGCU	124	AGCCAGTG GGCTAGCTACAACGA GGTTTGCT	1826
553	CAAACCCAC A CUGGCUUC	125	GAAGCCAG GGCTAGCTACAACGA GTGGTTTG	1827
557	CCACACUG G CUCUACCA	126	TGTAGAAAG GGCTAGCTACAACGA CAGTGTGG	1828
563	UGGCUUCU A CAGCUGCA	127	TGCAGCTG GGCTAGCTACAACGA AGAAGCCA	1829
566	CUUCUACAA G CUGCAAAU	128	ATTTGCAG GGCTAGCTACAACGA TGTAGAAG	1830
569	CUACAGCU G CAAAUUAUC	129	GATATTTG GGCTAGCTACAACGA AGCTGTAG	1831
573	AGCUGCAA A UAUCUAGC	130	GCTAGATA GGCTAGCTACAACGA TTGCAGCT	1832
575	CUGCAAAU A UCUAGCUG	131	CAGCTAGA GGCTAGCTACAACGA ATTTGCAG	1833
580	AAUAUCA G CUGUACCU	132	AGGTACAG GGCTAGCTACAACGA TAGATATT	1834
583	AUCUAGCU G UACCUACU	133	AGTAGGTA GGCTAGCTACAACGA AGCTAGAT	1835
585	CUAGCUGU A CCUACUUC	134	GAAGTAGG GGCTAGCTACAACGA ACAGCTAG	1836
589	CUGUACCU A CUUCAAAG	135	CTTTGAAG GGCTAGCTACAACGA AGGTACAG	1837
607	AGAAGGAA A CAGAAUCU	136	AGATTCTG GGCTAGCTACAACGA TTCCCTCT	1838
612	GAAACAGA A UCUGCAAU	137	ATTGCAGA GGCTAGCTACAACGA TCTGTTTC	1839
616	CAGAAUCU G CAAUCUAAU	138	ATAGATTG GGCTAGCTACAACGA AGATTCTG	1840
619	AAUCUGCA A UCUAUAAU	139	TATATAGA GGCTAGCTACAACGA TGCAGATT	1841
623	UGCAAUCA U UAUUAAA	140	TAAATATA GGCTAGCTACAACGA AGATTGCA	1842
625	CAAUCUAA U UAUUAAA	141	AATAAATA GGCTAGCTACAACGA ATAGATTG	1843
627	AUCUAUAA U UUUUAAA	142	CTAATAAA GGCTAGCTACAACGA ATATAGAT	1844
631	AUAUUAAA U UUAGUGAU	143	ATCACTAA GGCTAGCTACAACGA AAATATAT	1845
635	AUUUAAA G UGAAACAG	144	CTGTATCA GGCTAGCTACAACGA TAATAAAT	1846
638	UAUUAAGUG A UTACAGGU	145	TACCTGTA GGCTAGCTACAACGA CACTAATA	1847
640	UUAGUGAU A CAGGUAGA	146	TCTACCTG GGCTAGCTACAACGA ATCACTAA	1848
644	UGAUACAG G UAGACCUU	147	AAGGTCTA GGCTAGCTACAACGA CTGTATCA	1849
648	ACAGGUAG A CCUUUCGU	148	ACGAAAGG GGCTAGCTACAACGA CTACCTGT	1850
655	GACCUUUC G UAGAGAUG	149	CATCTCTA GGCTAGCTACAACGA GAAAGGTC	1851
661	UCGUAGAG A UGUACAGU	150	ACTGTACA GGCTAGCTACAACGA CTCTACGA	1852
663	GUAGAGAU G UTACAGUGA	151	TCACTGTA GGCTAGCTACAACGA ATCTCTAC	1853
665	AGAGAAGU A CAGUGAAA	152	TTTCACTG GGCTAGCTACAACGA ACATCTCT	1854

668	GAUGUACA G UGAAAUC	153	GGATTTCA GGCTAGCTACAACGA TGTACATC	1855
673	ACAGUGAA A UCCCCGAA	154	TTCCGGGA GGCTAGCTACAACGA TTCACTGT	1856
682	UCCCCGAA A UUAUACAC	155	GTGTATAA GGCTAGCTACAACGA TTCCGGGA	1857
685	CGAAGUUU A UACACAU	156	CATGTGTA GGCTAGCTACAACGA AATTTCGG	1858
687	GAAAUUUA A CACAU	157	GTCATGTG GGCTAGCTACAACGA ATAATTTC	1859
689	AAUUAUAC A CAUGACUG	158	CAGTCATG GGCTAGCTACAACGA GTATAATT	1860
691	UUAUACAC A UGACUGAA	159	TTCAGTCA GGCTAGCTACAACGA GTGTATAA	1861
694	UACACAU A CUGAAGGA	160	TCCCTPCAG GGCTAGCTACAACGA CATGTGTA	1862
708	GGAAGGGG G CUCGUCAU	161	ATGACGAG GGCTAGCTACAACGA TCCCTTCC	1863
712	GGGAGCUC G UCAUUCCG	162	GGGAATGA GGCTAGCTACAACGA GAGCTCCC	1864
715	AGCUCCUC A UUCCUCG	163	GCAGGGAA GGCTAGCTACAACGA GACGAGCT	1865
722	CAUUCCU G CCGGGUUA	164	TAACCCGG GGCTAGCTACAACGA AGGGAAATG	1866
727	CCUGCCGG G UUACGUCA	165	TGACGTAA GGCTAGCTACAACGA CCGGCAGG	1867
730	GCCGGGUU A CGUCACCU	166	AGGTGACG GGCTAGCTACAACGA AACCCGGC	1868
732	CGGGGUAC G UCACCUAA	167	TTAGGTGA GGCTAGCTACAACGA GTAACCCG	1869
735	GUUACGUC A CCUAACAU	168	ATGTTAGG GGCTAGCTACAACGA GACGTAAC	1870
740	GUCACCBA A CAUCACUG	169	CAGTGATG GGCTAGCTACAACGA TAGGTGAC	1871
742	CACCUAAC A UCACUGUU	170	AACAGTGA GGCTAGCTACAACGA GTTAGGTG	1872
745	CUAACAU A CUGUUACU	171	AGTAACAG GGCTAGCTACAACGA GATGTTAG	1873
748	ACAUCACU G UUACUUUA	172	TAAAGTAA GGCTAGCTACAACGA AGTGATGT	1874
751	UCACUGUU A CUUUAAAA	173	TTTTAAAG GGCTAGCTACAACGA AACAGTGA	1875
762	UUAAAAAA G UUUCACU	174	AGTGGAAA GGCTAGCTACAACGA TTTTTTAA	1876
768	AAGUUUCC A CUUGACAC	175	GTGTCAAG GGCTAGCTACAACGA GGAAACTT	1877
773	UCCACUUG A CACUUUGA	176	TCAAAGTG GGCTAGCTACAACGA CAAGTGG	1878
775	CACUUGAC A CUUUGAUC	177	GATCAAAG GGCTAGCTACAACGA GTCAAGTG	1879
781	ACACUUUG A UCCCCGAU	178	ATCAGGGAA GGCTAGCTACAACGA CAAAGTGT	1880
788	GAUCCUG A UGGAAAAC	179	TTTTTCCA GGCTAGCTACAACGA CAGGGATC	1881
795	GAUGGAAA A CGCAUAAU	180	ATTATGCG GGCTAGCTACAACGA TTTCCATC	1882
797	UGGAAAAC G CAUAAUCU	181	AGATTATG GGCTAGCTACAACGA TTTTTCCA	1883
799	GAAAACGC A UAAUCUGG	182	CCAGATTAA GGCTAGCTACAACGA GCGTTTTC	1884
802	AACGCAUA A UCUGGGAC	183	GTCCCAGA GGCTAGCTACAACGA TATGCGTT	1885
809	AAUCUGGG A CAGUAGAA	184	TTCTACTG GGCTAGCTACAACGA CCCAGATT	1886
812	CUGGGACA G UAGAAAGG	185	CCTTTCTA GGCTAGCTACAACGA TGTCCCAG	1887
821	UAGAAAGG G CUUCAUCA	186	TGATGAAG GGCTAGCTACAACGA CCTTTCTA	1888
826	AGGGCUUC A UCAUAUCA	187	TGATATGTA GGCTAGCTACAACGA GAAGCCCT	1889
829	GCUUCAUC A UAUAAA	188	ATTTGATA GGCTAGCTACAACGA GATGAAGC	1890
831	UUCAUCAU A UCAAAUGC	189	GCATTTGTA GGCTAGCTACAACGA ATGATGAA	1891
836	CAUAUCAA A UGCAACGU	190	ACGTTGCA GGCTAGCTACAACGA TTGATATG	1892
838	UAUAAA G CAACGUAC	191	GTACGTTG GGCTAGCTACAACGA ATTTGATA	1893
841	CAAAUGCA A CGUACAAA	192	TTTGTACG GGCTAGCTACAACGA TGCAATTG	1894
843	AAGGCAAC G UACAAAGA	193	TCTTTGTA GGCTAGCTACAACGA GTTGCATT	1895
845	UGCAACGU A CAAAGAAA	194	TTTCCTTGT GGCTAGCTACAACGA ACGTTGCA	1896
853	ACAAAGAA A UAGGGCUU	195	AAGCCCTA GGCTAGCTACAACGA TTCTTTGT	1897
858	GAAAIIAGG G CUUCUGAC	196	GTCAGAAAG GGCTAGCTACAACGA CCTATTTC	1898
865	GGCUUCUG A CCUGUGAA	197	TTCACAGG GGCTAGCTACAACGA CAGAAGCC	1899
869	UCUGACCU G UGAAGCAA	198	TTGCTTCA GGCTAGCTACAACGA AGGTCAGA	1900
874	CCUGUGAA G CAACAGUC	199	GACTGTTG GGCTAGCTACAACGA TTCACAGG	1901
877	GUGAAGCA A CAGUCAAU	200	ATTGACTG GGCTAGCTACAACGA TGCTTCAC	1902
880	AAGCAACA G UCAAUGGG	201	CCCATTGA GGCTAGCTACAACGA TGTTGCTT	1903
884	AACAGUCA A UGGGCCAU	202	AATGCCCA GGCTAGCTACAACGA TGACTGTT	1904
888	GUCAAUAGG G CAUJUUGUA	203	TACAAATG GGCTAGCTACAACGA CCATTGAC	1905
890	CAAUGGGC A UUUGUAIA	204	TATACAAA GGCTAGCTACAACGA GCCCATTG	1906

894	GGGCAUUU G UAUAAAGAC	205	GTCTTATA GGCTAGCTACAACGA AAATGCC	1907
896	GCAUUUGU A UAAGACAA	206	TTGCTTAA GGCTAGCTACAACGA ACAATGC	1908
901	UGUUAUAG A CAAACUAA	207	ATAGTTG GGCTAGCTACAACGA CTTATACA	1909
905	UAAGACAA A CUAUCUCA	208	TGAGATAG GGCTAGCTACAACGA TTGCTTA	1910
908	GACAAACU A UCUCACAC	209	GTGTGAGA GGCTAGCTACAACGA AGTTTGTC	1911
913	ACUAUCUC A CACAUCGA	210	TCGATGTG GGCTAGCTACAACGA GAGATAGT	1912
915	UAUCUCAC A CAUCGACA	211	TGTCGATG GGCTAGCTACAACGA GTGAGATA	1913
917	UCUCACAC A UCGACAAA	212	TTTGTGCGA GGCTAGCTACAACGA GTGTGAGA	1914
921	ACACAUCG A CAAACCAA	213	TTGGTTTG GGCTAGCTACAACGA CGATGTGT	1915
925	AUCGACAA A CCAAUACA	214	TGTATTGG GGCTAGCTACAACGA TTGTCGAT	1916
929	ACAAACCR A UACAAUCA	215	TGATTGTA GGCTAGCTACAACGA TGTTTGT	1917
931	AAACCAAU A CAAUCAUA	216	TATGATTG GGCTAGCTACAACGA ATGGTTT	1918
934	CCAAUACA A UCAUAGAU	217	ATCTATGA GGCTAGCTACAACGA TGTATTGG	1919
937	AUACAAUC A UAGAUGUC	218	GACATCTA GGCTAGCTACAACGA GATTGTAT	1920
941	AAUCAUAG A UGUCCAAA	219	TTTGGACA GGCTAGCTACAACGA CTATGATT	1921
943	UCAUAGAU G UCCAAAUA	220	TATTTGGA GGCTAGCTACAACGA ATCTATGA	1922
949	AUGUCCAA A UAAGCACA	221	TGTCTTAA GGCTAGCTACAACGA TTGGACAT	1923
953	CCAAUUA G CACACCAC	222	GTGGTGTG GGCTAGCTACAACGA TTATTGG	1924
955	AAAUUAGC A CACCACGC	223	GGCTGGTG GGCTAGCTACAACGA GCTTATT	1925
957	AUAAGCAC A CCACGCC	224	GGGCGTGG GGCTAGCTACAACGA GTGCTTAT	1926
960	AGCACACCC A CGCCCAGU	225	ACTGGGGC GGCTAGCTACAACGA GGTGTGCT	1927
962	CACACCAC G CCCAGUCA	226	TGACTGGG GGCTAGCTACAACGA GTGGTGTG	1928
967	CAAGCCCA G UCAAAUUA	227	TAATTTGA GGCTAGCTACAACGA TGGCGTG	1929
972	CCAGUCAA A UUACUUAG	228	CTAAGTAA GGCTAGCTACAACGA TTGACTGG	1930
975	GUCAAAUJ A CUUAGAGG	229	CCTCTAAG GGCTAGCTACAACGA AATTGAC	1931
983	ACUUAGAG G CCAUACUC	230	GAGTATGG GGCTAGCTACAACGA CTCTAAGT	1932
986	UAGAGGCC A UACUCUUG	231	CAAGAGTA GGCTAGCTACAACGA GCCCTCTA	1933
988	GAGGCCAU A CUCUUGUC	232	GACAAGAG GGCTAGCTACAACGA ATGGCCTC	1934
994	AUACUCUU G UCCUCAAU	233	ATTGAGGA GGCTAGCTACAACGA AAGAGTAT	1935
1001	UGUCCUCA A UUGUACUG	234	CACTACAA GGCTAGCTACAACGA TGAGGACA	1936
1004	CCUCAAUU G UACUGCUA	235	TAGCAGTA GGCTAGCTACAACGA AATTGAGG	1937
1006	UCAAUUGU A CUGCUACC	236	GGTAGCAG GGCTAGCTACAACGA ACAATTGA	1938
1009	AIUUGUACU G CUACCACU	237	AGTGGTAG GGCTAGCTACAACGA AGTACAAT	1939
1012	GUACUGCU A CCACUCCC	238	GGGAGTGG GGCTAGCTACAACGA AGCAGTAC	1940
1015	CUGCUACC A CUCCCCUUG	239	CAAGGGAG GGCTAGCTACAACGA GGTAGCAG	1941
1025	UCCCCUUGA A CACGAGAG	240	CTCTCGTG GGCTAGCTACAACGA TCAAGGGA	1942
1027	CCUUGAAC A CGAGAGUU	241	AACTCTCG GGCTAGCTACAACGA GTTCAAGG	1943
1033	ACACGGAGA G UUCAAAUG	242	CATTTGAA GGCTAGCTACAACGA TCTCGTGT	1944
1039	GAGUUCAA A UGACCUUG	243	CCAGGTCA GGCTAGCTACAACGA TTGAACTC	1945
1042	UUCAAAUG A CCUGGAGU	244	ACTCCAGG GGCTAGCTACAACGA CATTGAA	1946
1049	GACCGGGA G UUACCCUG	245	CAGGGTAA GGCTAGCTACAACGA TCCAGGTC	1947
1052	CUGGAGUU A CCCUGAUG	246	CATCAGGG GGCTAGCTACAACGA AACTCCAG	1948
1058	UUACCCUG A UGAAAAAA	247	TTTTTTCA GGCTAGCTACAACGA CAGGGTAA	1949
1067	UGAAAAAAA A UAAGAGAG	248	CTCTCTTA GGCTAGCTACAACGA TTTTTTCA	1950
1075	AUAAGAGA G CUUCCGUA	249	TACGGAAG GGCTAGCTACAACGA TCTCTTAT	1951
1081	GAGCUUCC G UAAGGCGA	250	TCGCCTTA GGCTAGCTACAACGA GGAAGCTC	1952
1086	UCCGUAAG G CGACGAUJ	251	ATTCGTCG GGCTAGCTACAACGA CTTACGGA	1953
1089	GUAAGGCG A CGAAUJUG	252	TCAATTCG GGCTAGCTACAACGA CGCCTTAC	1954
1093	GGCGACGA A UUGACCAA	253	TTGGTCAA GGCTAGCTACAACGA TCGTCGCC	1955
1097	ACGAUADUG A CCAAAGCA	254	TGCTTTGG GGCTAGCTACAACGA CAATTGCT	1956
1103	UGACCAAA G CAAUUCCC	255	GGGAATTG GGCTAGCTACAACGA TTTGGTCA	1957
1106	CCAAAGCA A UUCCCAUG	256	CATGGGAA GGCTAGCTACAACGA TGCTTTGG	1958

1112	CAAUUCCC A UGCCAACA	257	TGTTGGCA GGCTAGCTACAACGA GGGATTG	1959
1114	AUUCCCAU G CCAACAU	258	TATGTTGG GGCTAGCTACAACGA ATGGGAAT	1960
1118	CCAUCCA A CAUAIUUCU	259	AGAATATG GGCTAGCTACAACGA TGGCATGG	1961
1120	AUGCCAAC A UAUUCUAC	260	GTTAGATAA GGCTAGCTACAACGA GTTGGCAT	1962
1122	GCCAACAU A UUCUACAG	261	CTGTAGAA GGCTAGCTACAACGA ATGTTGGC	1963
1127	CAUAUUCU A CAGUGUUC	262	GAACACTG GGCTAGCTACAACGA AGAATATG	1964
1130	AUUCUAC A G UGUUCUUA	263	TAAGAAC A GGCTAGCTACAACGA TGTAGAAT	1965
1132	UCUACAGU G UUCUUACU	264	AGTAAGAA GGCTAGCTACAACGA ACTGTAGA	1966
1138	GUGUUCUU A CUAUUGAC	265	GTCAATAG GGCTAGCTACAACGA AAGAACAC	1967
1141	UUCUUACU A UUGACAAA	266	TTTGTCAA GGCTAGCTACAACGA AGTAAGAA	1968
1145	UACUAIUUG A CAAAUGC	267	GCATTTTG GGCTAGCTACAACGA CAATAGTA	1969
1150	UUGACAAA A UGGAGAAC	268	TTCTCTGA GGCTAGCTACAACGA TTTGTCAA	1970
1152	GACAAAAU G CAGAACAA	269	TTGTTCTG GGCTAGCTACAACGA ATTTTGTC	1971
1157	AAUGCAGA A CAAAGACA	270	TGTCTTTG GGCTAGCTACAACGA TCTGCATT	1972
1163	GAACAAAG A CAAAGGAC	271	GTCCTTTG GGCTAGCTACAACGA CTTTGTTC	1973
1170	GACAAAGG A CUUUUAC	272	GTATAAAG GGCTAGCTACAACGA CCTTTGTC	1974
1175	AGGACUUU A UACUUGUC	273	GACAAGTA GGCTAGCTACAACGA AAAGTCCT	1975
1177	GACUUIAU A CUUGUCGU	274	ACGACAAG GGCTAGCTACAACGA ATAAAGTC	1976
1181	UUAUACUU G UCGUGUAA	275	TTACACGA GGCTAGCTACAACGA AAGTATAA	1977
1184	UACUUGUC G UGUAAGGA	276	TCCTTACA GGCTAGCTACAACGA GACAAGTA	1978
1186	CUUGUCGU G UAAGGAGU	277	ACTCCTTA GGCTAGCTACAACGA ACGACAAG	1979
1193	UGUAAGGA G UGGACCAU	278	ATGGTCCA GGCTAGCTACAACGA TCCTTACA	1980
1197	AGGAGUGG A CCAUCAU	279	AATGATGG GGCTAGCTACAACGA CCACTCCT	1981
1200	AGUGGACC A UCAUUCAA	280	TTGAATGA GGCTAGCTACAACGA GGTCCACT	1982
1203	GGACCAUC A UUAAAUC	281	GATTTGAA GGCTAGCTACAACGA GATGGTCC	1983
1209	UCAUUCAA A UCUGUJAA	282	TTAACAGA GGCTAGCTACAACGA TTGAATGA	1984
1213	UCAAAUCU G UUAACACC	283	GGTGTAA GGCTAGCTACAACGA AGATTTGA	1985
1217	AUCUGUUA A CACCUCAG	284	CTGAGGTG GGCTAGCTACAACGA TAACAGAT	1986
1219	CUGUUAAC A CCUCAGUG	285	CACTGAGG GGCTAGCTACAACGA TTAAACAG	1987
1225	ACACCUCA G UGCAUATA	286	TATATGCA GGCTAGCTACAACGA TGAGGTGT	1988
1227	ACCUAGU G CAUAUJUA	287	TATATATG GGCTAGCTACAACGA ACTGAGGT	1989
1229	CUCAGUGC A UAUUAUAG	288	CATATATA GGCTAGCTACAACGA GCACTGAG	1990
1231	CAGUGCAU A UAU AUGAU	289	ATCATATA GGCTAGCTACAACGA ATGCACTG	1991
1233	GGCAUUAU A UAU GUAUA	290	TTATCATA GGCTAGCTACAACGA ATATGCAC	1992
1235	GCAUUAU A UGAUAAAG	291	CTTTATCA GGCTAGCTACAACGA ATATATGC	1993
1238	UAAUUAUAG A UAAAGCAU	292	ATGCTTTA GGCTAGCTACAACGA CATATATA	1994
1243	AUGAUAAA G CAIJUCAUC	293	GATGAATG GGCTAGCTACAACGA TTTATCAT	1995
1245	GAUAAAAGC A UUCAUCAC	294	GTGATGAA GGCTAGCTACAACGA GCTTTATC	1996
1249	AAGCAUUC A UCACUGUG	295	CACAGTGA GGCTAGCTACAACGA GAATGCTT	1997
1252	CAUUCAUC A CUGUGAAA	296	TTTCACAG GGCTAGCTACAACGA GATGAATG	1998
1255	UCAUCACU G UGAACCAU	297	ATGTTTCA GGCTAGCTACAACGA AGTGTATGA	1999
1260	ACUGUGAA A CAUCGAAA	298	TTTCGATG GGCTAGCTACAACGA TTTCACAGT	2000
1262	UGUGAAC A UCGAAAAC	299	GTTCGATG GGCTAGCTACAACGA GTTTCACA	2001
1269	CAUCGAAA A CAGCAGGU	300	ACCTGCTG GGCTAGCTACAACGA TTTCGATG	2002
1272	CGAAAACA G CAGGUGCU	301	AGCACCTG GGCTAGCTACAACGA TGTTTTCG	2003
1276	AAACGGAG G UGCUUGAA	302	TTCAAGCA GGCTAGCTACAACGA CTGCTGTT	2004
1278	CAGCAGGU G CUUGAAAC	303	GTTCAGCA GGCTAGCTACAACGA ACCTGCTG	2005
1285	UGCUUGAA A CCCUAGCU	304	AGCTACGG GGCTAGCTACAACGA TTCAAGCA	2006
1288	UUGAAACC G UAGCUGGC	305	GCCAGCTA GGCTAGCTACAACGA GGTTTCAA	2007
1291	AAACCGUA G CUGGCAAG	306	CTTGCCAG GGCTAGCTACAACGA TACGGTTT	2008
1295	CGUAGCUG G CAAGCGGU	307	ACCGCTTG GGCTAGCTACAACGA CAGCTACG	2009
1299	GCUGGCAA G CGGUCUUA	308	TAAGACCG GGCTAGCTACAACGA TTGCCCCG	2010

1302	GGCAAGCG G UCUUACCG	309	CGGTAAGA GGCTAGCTACAACGA CGCTTGCC	2011
1307	CGGGUCUU A CCGGCUCU	310	AGAGCCGG GGCTAGCTACAACGA AAGACCGC	2012
1311	UCUUACCG G CUCUCUAU	311	ATAGAGAG GGCTAGCTACAACGA CGGTAAGA	2013
1318	GCGCUCUCU A UGAAAGUG	312	CACTTTCA GGCTAGCTACAACGA AGAGAGCC	2014
1324	CUAUGAAA G UGAAGGCA	313	TGCCCTCA GGCTAGCTACAACGA TTTCATAG	2015
1330	AAGUGAAG G CAUUCUCC	314	GGGAAATG GGCTAGCTACAACGA CTTCACTT	2016
1332	GUGAAGGC A UUUCCCUC	315	GAGGGAAA GGCTAGCTACAACGA GCCTTCAC	2017
1341	UUUCCCUC G CCGGAAGU	316	ACTTCCGG GGCTAGCTACAACGA GAGGGAAA	2018
1348	CGCCGGAA G UUGUUAUGG	317	CCATACAA GGCTAGCTACAACGA TTCCGGCG	2019
1351	CGGAAGUU G UAUGGUUA	318	TAACCATA GGCTAGCTACAACGA AACTTCGG	2020
1353	GAAGUUGU A UGGUUTAA	319	TTTAACCA GGCTAGCTACAACGA ACAACTTC	2021
1356	GUUGUAUG G UAAAAAGA	320	TCTTTAA GGCTAGCTACAACGA CATAAAC	2022
1364	GUAAAAG A UGGGUUAC	321	GTAACCCA GGCTAGCTACAACGA CTTTTAAC	2023
1368	AAAGAUGG G UUACCUGC	322	GCAGGTAA GGCTAGCTACAACGA CCATCTTT	2024
1371	GAUGGGUU A CCUGCGAC	323	GTCGCAGG GGCTAGCTACAACGA AACCCATC	2025
1375	GGUUACCU G CGACUGAG	324	CTCAGTCG GGCTAGCTACAACGA AGGTAACC	2026
1378	UACCUUGCG A CUGAGAAA	325	TTTCTCAG GGCTAGCTACAACGA CGCAGGTA	2027
1386	ACUAGAGAA A UCUGCUCG	326	CGAGCAGA GGCTAGCTACAACGA TTCTCAGT	2028
1390	AGAAAUCU G CUCGCUAU	327	ATAGCGAG GGCTAGCTACAACGA AGATTTCT	2029
1394	AUCUGCUC G CUAUUUGA	328	TCAAATAG GGCTAGCTACAACGA GAGCAGAT	2030
1397	UGCUCGCU A UUUGACUC	329	GAGTCAAA GGCTAGCTACAACGA AGCGAGCA	2031
1402	CCAUUUUG A CUCGUGGC	330	GCCACGAG GGCTAGCTACAACGA CAAATAGC	2032
1406	UUUGACUC G UGGCUACU	331	AGTAGCCA GGCTAGCTACAACGA GAGTCAAA	2033
1409	GACUCGUG G CUACUCGU	332	ACGAGTAG GGCTAGCTACAACGA CACGAGTC	2034
1412	UCGUGGGCU A CUCGUUAA	333	TTAACGAG GGCTAGCTACAACGA AGCCACGA	2035
1416	GGCUACUC G UJAAUUAU	334	ATAATTAA GGCTAGCTACAACGA GAGTAGCC	2036
1420	ACUCGUUA A UUAUCAAG	335	CTTGATAA GGCTAGCTACAACGA TAACGAGT	2037
1423	CGUAAAUU A UCAAGGAC	336	GTCCTTGA GGCTAGCTACAACGA AATTAACG	2038
1430	UAUCAAGG A CGUAAACUG	337	CAGTTACG GGCTAGCTACAACGA CCTTGATA	2039
1432	UCAAGGAC G UUACUGAA	338	TTCAGTTA GGCTAGCTACAACGA GTCCCTTGA	2040
1435	AGGACGUA A CUGAAGAG	339	CTCTTCAG GGCTAGCTACAACGA TACGTCCT	2041
1445	UGAAGAGG A UGCAGGGG	340	TCCCTGCA GGCTAGCTACAACGA CCTCTTCA	2042
1447	AAGAGGAU G CAGGGAAU	341	ATTCCTG GGCTAGCTACAACGA ATCCTCTT	2043
1454	UGCAGGGG A UUAUACAA	342	TTGTATAA GGCTAGCTACAACGA TCCCTGCA	2044
1457	AGGGAAUU A UACAAUCU	343	AGATTGTA GGCTAGCTACAACGA AATTCCT	2045
1459	GGAAUUAU A CAAUCUUG	344	CAAGATTG GGCTAGCTACAACGA ATAATTCC	2046
1462	AUUAUACAA A UCUUGCUG	345	CAGCAAGA GGCTAGCTACAACGA TGTATAAT	2047
1467	ACAAUCUU G CUGAGCAU	346	ATGCTCAG GGCTAGCTACAACGA AAGATTGT	2048
1472	CUUGCUGA G CAUAAAAC	347	GTTTTATG GGCTAGCTACAACGA TCAGCAAG	2049
1474	UGCUGAGC A UAAAACAG	348	CTGTTTTA GGCTAGCTACAACGA GCTCAGCA	2050
1479	ACCAUAAA A CAGUCAAA	349	TTTGACTG GGCTAGCTACAACGA TTTATGCT	2051
1482	AUAAAACA G UCAAAUGU	350	ACATTGTA GGCTAGCTACAACGA TGTTTTAT	2052
1487	ACAGUCAA A UGUGUUUA	351	TAAACACA GGCTAGCTACAACGA TTGACTGT	2053
1489	AGUCAAAU G UGUUUAAA	352	TTTAAACA GGCTAGCTACAACGA ATTTGACT	2054
1491	UCAAAUGU G UUUAAAAA	353	TTTTTAAA GGCTAGCTACAACGA ACATTGTA	2055
1499	GUUUAAAA A CCUCACUG	354	CAGTGAGG GGCTAGCTACAACGA TTTTAAAC	2056
1504	AAAACCUC A CUGCCACU	355	AGTGGCAG GGCTAGCTACAACGA GAGGTTTT	2057
1507	ACCUCACU G CCACUCUA	356	TAGAGTGG GGCTAGCTACAACGA AGTGAGGT	2058
1510	UCACUGCC A CUCUAAU	357	AATTAGAG GGCTAGCTACAACGA GGCAGTGA	2059
1516	CCACUCUA A UUGUCAAU	358	ATTGACAA GGCTAGCTACAACGA TAGAGTGG	2060
1519	CUCUAAU G UCAAUGUG	359	CACATTGA GGCTAGCTACAACGA AATTAGAG	2061
1523	AAUUGUCA A UGUGAAC	360	GTTCACCA GGCTAGCTACAACGA TGACAAATT	2062

1525	UUGUCAAU G UGAAACCC	361	GGGTTTCA GGCTAGCTACAACGA ATTGACAA	2063
1530	AAUGUGAA A CCCCAGAU	362	ATCTGGGG GGCTAGCTACAACGA TTCACATT	2064
1537	AACCCCCAG A UUUACGAA	363	TTCGTAAA GGCTAGCTACAACGA CTGGGGTT	2065
1541	CCAGAUUU A CGAAAAGG	364	CCTTTTCG GGCTAGCTACAACGA AAATCTGG	2066
1549	ACGAAAAG G CCGUGUCA	365	TGACACGG GGCTAGCTACAACGA CTTTTCGT	2067
1552	AAAAGGCC G UGUCAUCG	366	CGATGACA GGCTAGCTACAACGA GGCCCTTT	2068
1554	AAGGCCGU G UCAUCGUU	367	AACGATGA GGCTAGCTACAACGA ACGGCCTT	2069
1557	GCCGUGUC A UCGUUUCC	368	GGAAACCA GGCTAGCTACAACGA GACACGGC	2070
1560	GUGUCAUC G UUUCAGA	369	TCTGGAAA GGCTAGCTACAACGA GATGACAC	2071
1568	GUUUCAG A CCCGGCUC	370	GAGCCGGG GGCTAGCTACAACGA CTGGAAAC	2072
1573	CAGACCCC G CUCUCUAC	371	GTAGAGAG GGCTAGCTACAACGA CGGGCTCG	2073
1580	GGCUCUCU A CCCACUGG	372	CCAGTGGG GGCTAGCTACAACGA AGAGAGCC	2074
1584	CUCUACCC A CUGGGCAG	373	CTGCCCG AG GGCTAGCTACAACGA GGGTAGAG	2075
1589	CCCACUGG G CAGCAGAC	374	GTCTGCTG GGCTAGCTACAACGA CCAGTGGG	2076
1592	ACUGGGCA G CAGACAAA	375	TTTGTCTG GGCTAGCTACAACGA TGCCCGAT	2077
1596	GGCAGCGA G CAAAUCCU	376	AGGATTTG GGCTAGCTACAACGA CTGCTGCC	2078
1600	GCAGACAA A UCCUGACU	377	AGTCAGGA GGCTAGCTACAACGA TTGTCTGC	2079
1606	AAAUCUG A CUUGUACC	378	GGTACAAAG GGCTAGCTACAACGA CAGGATTT	2080
1610	CCUGACUU G UACCGCAU	379	ATGCGGTA GGCTAGCTACAACGA AAGTCAGG	2081
1612	UGACUUGU A CCGCAUUA	380	ATATGCGG GGCTAGCTACAACGA ACAAGTCA	2082
1615	CUUGUACC G CAUAAUGG	381	ACCATATG GGCTAGCTACAACGA GGTACAAG	2083
1617	UGUACCGC A UAUGGUAU	382	ATACCATA GGCTAGCTACAACGA GCGGTACA	2084
1619	UACCGCAU A UGGUAUCC	383	GGATACCA GGCTAGCTACAACGA ATGCGGTA	2085
1622	CGCAUAAUG G UAUCCCCU	384	GAGGGATA GGCTAGCTACAACGA CATATGCG	2086
1624	CAUAAUGG U UCCCCUAA	385	TTGAGGGG GGCTAGCTACAACGA ACCATATG	2087
1632	AUCCCCUCA A CCUACAAU	386	ATTGTAGG GGCTAGCTACAACGA TGAGGGAT	2088
1636	CUCAACCU A CAAUCAAG	387	CTTGATIG GGCTAGCTACAACGA AGGTTGAG	2089
1639	AACCUACA A UCAAGUGG	388	CCACTTGA GGCTAGCTACAACGA TGTAGGTT	2090
1644	ACAAUCAA G UGGUUCUG	389	CAGAACCA GGCTAGCTACAACGA TTGATTGT	2091
1647	AUCAAGUG G UUCUGGCA	390	TGCCAGAA GGCTAGCTACAACGA CACTTGAT	2092
1653	UGGUUCUG G CACCCUG	391	CAGGGITG GGCTAGCTACAACGA CAGAACCA	2093
1655	GUUCUGGC A CCCCUGUA	392	TACAGGGG GGCTAGCTACAACGA GCCAGAAC	2094
1661	GCACCCCU G UAACCAAU	393	TATGGTTA GGCTAGCTACAACGA AGGGGTGC	2095
1664	CCCCUGUA A CCAUAAUC	394	GATTATGG GGCTAGCTACAACGA TACAGGGG	2096
1667	CUGUAACC A UAAUCAUU	395	AATGATTA GGCTAGCTACAACGA GGTTACAG	2097
1670	UAACCAAUA A UCAUUCCG	396	CGGAATGA GGCTAGCTACAACGA TATGGTTA	2098
1673	CCAUAAUC A UUCCGAAG	397	CTTCGGAA GGCTAGCTACAACGA GATTATGG	2099
1681	AUUCGAA G CAAGGUGU	398	ACACCTTG GGCTAGCTACAACGA TTGGAAAT	2100
1686	GAAGCAAG G UGUGACUU	399	AAGTCACA GGCTAGCTACAACGA CTTGCTTC	2101
1688	AGCAAGGU G UGACUUUU	400	AAAAGTCA GGCTAGCTACAACGA ACCTTGCT	2102
1691	AAGGUGUG A CUUUGGU	401	AAACAAAG GGCTAGCTACAACGA CACACCTT	2103
1697	UGACUUUU G UUCCAAUA	402	TATTGGAA GGCTAGCTACAACGA AAAAGTCA	2104
1703	UUGUUCCA A UAAUGAAG	403	CTTCATTGA GGCTAGCTACAACGA TGGAACAA	2105
1706	UUCCAAUA A UGAAGAGU	404	ACTCTTCA GGCTAGCTACAACGA TATTGGAA	2106
1713	AAUGAAGA G UCCUUUUAU	405	ATAAAAGG GGCTAGCTACAACGA TCTTCATT	2107
1720	AGUCCUUU A UCCUGGAU	406	ATCCAGGA GGCTAGCTACAACGA AAAGGACT	2108
1727	UAUCCUGG A UGCUGACA	407	TGTCAAGCA GGCTAGCTACAACGA CCAGGATA	2109
1729	UCCUGGAU G CUGACAGC	408	GCTGTCAG GGCTAGCTACAACGA ATCCAGGA	2110
1733	GGAUGGUG A CAGCAACA	409	TGTTGCTG GGCTAGCTACAACGA CAGCATCC	2111
1736	UGCUGACA G CAACAUGG	410	CCATGTTG GGCTAGCTACAACGA TGTCAAGCA	2112
1739	UGACAGCA A CAUGGGAA	411	TTCCCATG GGCTAGCTACAACGA TGCTGTCA	2113
1741	ACAGCAAC A UGGGAAAC	412	GTTCCTCA GGCTAGCTACAACGA GTTGCTGT	2114

1748	CAUGGGAA A CAGAAUUG	413	CAATTCTG GGCTAGCTACAACGA TTCCCATG	2115
1753	GAAACAGA A UUGAGAGC	414	GCTCTCAA GGCTAGCTACAACGA TCTGTTTC	2116
1760	AAUUGAGA G CAUCACUC	415	GAGTGATG GGCTAGCTACAACGA TCTCAATT	2117
1762	UUGAGAGC A UCACUCAG	416	CTGAGTGA GGCTAGCTACAACGA GCTCTCAA	2118
1765	AGAGCAUC A CUCAGCGC	417	GCGCTGAG GGCTAGCTACAACGA GATGCTCT	2119
1770	AUCACUCA G CGCAUGGC	418	GCCATGCG GGCTAGCTACAACGA TGAGTGAT	2120
1772	CACUCAGC G CAUGGCAA	419	TTGCCATG GGCTAGCTACAACGA GCTGAGTG	2121
1774	CUCAGCGC A UGGCAATA	420	TATTGCCA GGCTAGCTACAACGA GCGCTGAG	2122
1777	AGCGCAUG G CAAUAUUA	421	TATTATTG GGCTAGCTACAACGA CATGCGCT	2123
1780	GCAUGGCA A UAAUAGAA	422	TTCTATTA GGCTAGCTACAACGA TGCCATGC	2124
1783	UGGCAUA A UAGAAGGA	423	TCCCTCTA GGCTAGCTACAACGA TATTGCCA	2125
1796	AGGAAAGA A UAAGAUGG	424	CCATCTTA GGCTAGCTACAACGA TCTTTCCCT	2126
1801	AGAAUAG A UGGCUAGC	425	GCTAGCCA GGCTAGCTACAACGA CTTATTCT	2127
1804	AUAAGAUG G CUAGCACC	426	GGTGCTAG GGCTAGCTACAACGA CATCTTAT	2128
1808	GAUGGGCUA G CACCUUGG	427	CCAAGGTG GGCTAGCTACAACGA TAGCCATC	2129
1810	UGGCUAGC A CCUUGGUU	428	AACCAAGG GGCTAGCTACAACGA GCTAGCCA	2130
1816	GCACCUUG G UUGGUGGU	429	AGCCACAA GGCTAGCTACAACGA CAAGGTGC	2131
1819	CCUUGGUU G UGGCUGAC	430	GTCAGCCA GGCTAGCTACAACGA AACCAAGG	2132
1822	UGGUUGUG G CUGACUCU	431	AGAGTCAG GGCTAGCTACAACGA CACAACCA	2133
1826	UGUGGCUG A CUCUAGAA	432	TTCTAGAG GGCTAGCTACAACGA CAGCCACA	2134
1834	ACUCUAGA A UUUCCUGGA	433	TCCAGAAA GGCTAGCTACAACGA TCTAGAGT	2135
1843	UUUCUGGA A UCUACADU	434	AATGTAGA GGCTAGCTACAACGA TCCAGAAA	2136
1847	UGGAAUCU A CAUUGCA	435	TGCAAATG GGCTAGCTACAACGA AGATTCCA	2137
1849	GAAUCUAC A UUUGCAUA	436	TATGCAA GGCTAGCTACAACGA GTAGATTG	2138
1853	CUACAUUU G CAUAGCUU	437	AAGCTATG GGCTAGCTACAACGA AAATGTAG	2139
1855	ACAUUUGC A UAGCUUCC	438	GGAAAGCTA GGCTAGCTACAACGA GCAAATGT	2140
1858	UUUGCAUA G CUUCCAAU	439	ATTGGAAG GGCTAGCTACAACGA TATGCAA	2141
1865	AGCUUCCA A UAAAGUUG	440	CAACTTTA GGCTAGCTACAACGA TGGAGCT	2142
1870	CCAAUAAA G UGGGGACU	441	AGTCCCAA GGCTAGCTACAACGA TTTATTGG	2143
1876	AAGUUGGG A CUGUGGG	442	TCCCCACAG GGCTAGCTACAACGA CCCAACCT	2144
1879	UUGGGACU G UGGGAAGA	443	TCTTCCCA GGCTAGCTACAACGA AGTCCCAA	2145
1889	GGGAAGAA A CAUAAAGCU	444	AGCTTATG GGCTAGCTACAACGA TTCTTCCC	2146
1891	GAAGAAC A UAAGCUUU	445	AAAGCTTA GGCTAGCTACAACGA GTTCTTC	2147
1895	AAACAUAA G CUUUUAUA	446	TATAAAAG GGCTAGCTACAACGA TTATGTTT	2148
1901	AAGCUUUU A UAUCACAG	447	CTGTGATA GGCTAGCTACAACGA AAAAGCTT	2149
1903	GCUUUUUAU A UCACAGAU	448	ATCTGTGA GGCTAGCTACAACGA ATAAAAGC	2150
1906	UUUUAUAC A CAGAUGUG	449	CACATCTG GGCTAGCTACAACGA GATATAAA	2151
1910	UAUCACAG A UGUGCCAA	450	TTGGCACA GGCTAGCTACAACGA CTGTGATA	2152
1912	UCACAGAU G UGCCAAAU	451	ATTTGGCA GGCTAGCTACAACGA ATCTGTGA	2153
1914	ACAGAUGU G CCAAAUUG	452	CCATTTGG GGCTAGCTACAACGA ACATCTGT	2154
1919	UGUGCCAA A UGGGUUUC	453	GAACCCCA GGCTAGCTACAACGA TTGGCACA	2155
1923	CCAAUUGG G UUUCAUGU	454	ACATGAAA GGCTAGCTACAACGA CCATTTGG	2156
1928	UGGGUUUC A UGUUUAACU	455	AGTTAACCA GGCTAGCTACAACGA GAAACCCA	2157
1930	GGUUUCAU G UUAACUUG	456	CAAGTTAA GGCTAGCTACAACGA ATGAAACC	2158
1934	UCAUGUUA A CUUGGAAA	457	TTTCCAAG GGCTAGCTACAACGA TAACATGA	2159
1945	UGGAAAAA A UGCCGACG	458	CGTCGGCA GGCTAGCTACAACGA TTTTTCCA	2160
1947	AAAAAAAU G CCGACGGA	459	TCCGTCGG GGCTAGCTACAACGA ATTTTTTC	2161
1951	AAADGCCG A CGGAAGGA	460	TCCCTCCG GGCTAGCTACAACGA CGGCATTT	2162
1964	AGGAGAGG A CCUGAAC	461	GTTTCAGG GGCTAGCTACAACGA CCTCTCCCT	2163
1971	GACCUGAA A CUGUCUUG	462	CAAGACAG GGCTAGCTACAACGA TTCAGGTC	2164
1974	CUGAAACU G UCUUGCAC	463	GTGCAAGA GGCTAGCTACAACGA AGTTTCAG	2165
1979	ACUGUCUJU G CACAGUUA	464	TAACGTG GGCTAGCTACAACGA AAGACAGT	2166

1981	UGUCUUGC A CAGUUUAC	465	GTAACTG GGCTAGCTACAACGA GCAAGACA	2167
1984	CUUGCAC A UUAACAAG	466	CTTGTAA GGCTAGCTACAACGA TGTGCAAG	2168
1988	CACAGUUA A CAAGUUCU	467	AGAACTTG GGCTAGCTACAACGA TAACTGTG	2169
1992	GUUAACAA G UUCUUUUA	468	TATAAGAA GGCTAGCTACAACGA TTGTTAAC	2170
1998	AAGUUCUU A UACAGAGA	469	TCTCTGTA GGCTAGCTACAACGA AAGAACTT	2171
2000	GUUCUUAU A CAGAGACG	470	CGTCTCTG GGCTAGCTACAACGA ATAAGAAC	2172
2006	AUACAGAG A CGUUUACU	471	AAGTAACG GGCTAGCTACAACGA CTCTGTAT	2173
2008	ACAGAGAC G UUACUJUGG	472	CCAAGTAA GGCTAGCTACAACGA GTCTCTGT	2174
2011	GAGACGUU A CUUUGGAIU	473	AATCCAAG GGCTAGCTACAACGA AACGTCTC	2175
2017	UUACUUGG A UUUUACUG	474	CAGTAAA GGCTAGCTACAACGA CCAAGTAA	2176
2022	UGGAJUUU A CUGCGGAC	475	GTCCGGAG GGCTAGCTACAACGA AAAATCCA	2177
2025	AUUUUACU G CGGACAGU	476	ACTGTCCG GGCTAGCTACAACGA AGTAAAAT	2178
2029	UACUGCGG A CAGUUAU	477	ATTAACTG GGCTAGCTACAACGA CCGCAGTA	2179
2032	UGCGGACA G UUAAUAAAC	478	GTATTAA GGCTAGCTACAACGA TGTCCGCA	2180
2036	GACAGUUA A UAACAGAA	479	TTCTGTTA GGCTAGCTACAACGA TAACTGTC	2181
2039	AGUUAUUA A CAGAACAA	480	TTGTTCTG GGCTAGCTACAACGA TATTAACT	2182
2044	AUAACAGA A CAAUGCAC	481	GTGCATTG GGCTAGCTACAACGA TCTGTTAT	2183
2047	ACAGAAC A UGCACUAC	482	GTAGTGCA GGCTAGCTACAACGA TGTTCTGT	2184
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2054	AAUGCACU A CAGUUAU	485	TAATACTG GGCTAGCTACAACGA AGTGCATT	2187
2057	GCACUACA G UAUUAGCA	486	TGCTAATA GGCTAGCTACAACGA TGTAGTGC	2188
2059	ACUACAGU A UUTAGCAAG	487	CTTGTAA GGCTAGCTACAACGA ACTGTAGT	2189
2063	CAGUUAUA G CAAGCTAA	488	TTTGTCTG GGCTAGCTACAACGA TAATACTG	2190
2067	AUUAGCAA G CAAAAAAAU	489	ATTTTTTG GGCTAGCTACAACGA TTGCTAAT	2191
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2077	AAAAAAUG G CCAUCACU	491	AGTGATGG GGCTAGCTACAACGA CATTTTTT	2193
2080	AAAUGGCC A UCACUUAAG	492	CTTAGTGA GGCTAGCTACAACGA GGCCATT	2194
2083	UGGCCAUC A CUAAGGAG	493	CTCCCTTG GGCTAGCTACAACGA GATGGCCA	2195
2091	ACUAAGGA G CACUCCAU	494	ATGGAGTG GGCTAGCTACAACGA TCCCTAGT	2196
2093	UAAGGAGC A CUCCAUCA	495	TGATGGAG GGCTAGCTACAACGA GCTCCTTA	2197
2098	AGCACUCC A UCACUCUU	496	AAGAGTGA GGCTAGCTACAACGA GGAGTGCT	2198
2101	ACUCCAUC A CUCUUAU	497	ATTAAGAG GGCTAGCTACAACGA GATGGAGT	2199
2108	CACCUUA A UCUUACCA	498	TGGTAAGA GGCTAGCTACAACGA TAAGAGTG	2200
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2116	AUCUUACC A UCAUGAAU	500	ATTCATGTA GGCTAGCTACAACGA GGTAAGAT	2202
2119	UUACCAUC A UGAAUGUU	501	AACATTCA GGCTAGCTACAACGA GATGGTAA	2203
2123	CAUCUAUG A UGUUUUCCC	502	GGGAAACA GGCTAGCTACAACGA TCATGATG	2204
2125	UCAUGAAU G UUUCCCCUG	503	CAGGGAAA GGCTAGCTACAACGA ATTCACTGA	2205
2133	GUUUCCU G CAAGADUC	504	GAATCTTG GGCTAGCTACAACGA AGGGAAAC	2206
2138	CCUGCRAG A UUCAGGCC	505	TGCCCTGAA GGCTAGCTACAACGA CTTGCAGG	2207
2144	AGAUUCAG G CACCUCU	506	CATAGGTG GGCTAGCTACAACGA CTGAAATCT	2208
2146	AUUCAGGC A CCUAUGCC	507	GGCATAGG GGCTAGCTACAACGA GCCTGAAT	2209
2150	AGGCACCU A UGCCUGCA	508	TGCAGGCC GGCTAGCTACAACGA AGGTGCCT	2210
2152	GCACCUAU G CCUGCAGA	509	TCTGCAGG GGCTAGCTACAACGA ATAGGTGC	2211
2156	CUAUGCCU G CAGAGCCA	510	TGGCTCTG GGCTAGCTACAACGA AGGCATAG	2212
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2168	AGCCAGGA A UGUAIUACA	512	TGTATACA GGCTAGCTACAACGA TCCCTGGCT	2214
2170	CCAGGAAU G UAUACACA	513	TGTGTATA GGCTAGCTACAACGA ATTCCCTGG	2215
2172	AGGAADGU A UACACAGG	514	CCTGTGTA GGCTAGCTACAACGA ACATTCT	2216
2174	GAAUGUUA A CACAGGGG	515	CCCCCTGTG GGCTAGCTACAACGA ATACATTC	2217
2176	AUGUAIAC A CAGGGGAA	516	TTCCCCCTG GGCTAGCTACAACGA GTATACAT	2218

2188	GGGAAGAA A UCCUCCAG	517	CTGGAGGA GGCTAGCTACAACGA TTCTTCCC	2219
2206	AGAAAAGAA A UUACAAUC	518	GATTGTAA GGCTAGCTACAACGA TTCTTTCT	2220
2209	AAGAAAUU A CAAUCAGA	519	TCTGATTG GGCTAGCTACAACGA AATTTCTT	2221
2212	AAAUUACA A UCAGAGAU	520	ATCTCTGA GGCTAGCTACAACGA TGTAATTT	2222
2219	AAUCAGAG A UCAGGAAG	521	CTTCCTGA GGCTAGCTACAACGA CTCTGATT	2223
2227	AUCAGGAA G CACCAUAC	522	GTATGGTG GGCTAGCTACAACGA TTCCCTGAT	2224
2229	CAGGAAGC A CCAUACCU	523	AGGTATGG GGCTAGCTACAACGA GCTTCCTG	2225
2232	GAAGCACC A UACCUCCU	524	AGGAGGTA GGCTAGCTACAACGA GGTGCTTC	2226
2234	AGCACCAU A CCUCUGC	525	GCAGGAGG GGCTAGCTACAACGA ATGGTGCT	2227
2241	UACCUCCU G CGAAACCU	526	AGGTTTCG GGCTAGCTACAACGA AGGAGGTA	2228
2246	CCUGCGAA A CCUCAGUG	527	CACTGAGG GGCTAGCTACAACGA TTCCGAGG	2229
2252	AAACCUCA G UGAUCACA	528	TGTGATCA GGCTAGCTACAACGA TGAGGTTT	2230
2255	CCUCAGUG A UCACACAG	529	CTGTGTGA GGCTAGCTACAACGA CACTGAGG	2231
2258	CAGUGAUC A CACAGUGG	530	CCACTGTG GGCTAGCTACAACGA GATCACTG	2232
2260	GUGAUCAC A CAGUGGCC	531	GGCCACTG GGCTAGCTACAACGA GTGATCAC	2233
2263	AUCACACA G UGGCCAUC	532	GATGGCCA GGCTAGCTACAACGA TGTGTGAT	2234
2266	ACACAGUG G CCAUCAGC	533	GCTGATGG GGCTAGCTACAACGA CACTGTGT	2235
2269	CAGUGGCC A UCAGCAGU	534	ACTGCTGA GGCTAGCTACAACGA GGCCACTG	2236
2273	GGCCAUC A CAGUUCCA	535	TGGAACTG GGCTAGCTACAACGA TGATGGCC	2237
2276	CAUCAGCA G UUCCACCA	536	TGGTGGAA GGCTAGCTACAACGA TGCTGATG	2238
2281	GCAGUUCC A CCACUUUA	537	TAAAGTGG GGCTAGCTACAACGA GGAACCTGC	2239
2284	GUUCCACC A CUUUAGAC	538	GTCTAAAG GGCTAGCTACAACGA GGTGGAAC	2240
2291	CACIUUAG A CUGUCAUG	539	CATGACAG GGCTAGCTACAACGA CTAAACTG	2241
2294	UUUAGACU G UCAUGCUA	540	TAGCATGA GGCTAGCTACAACGA AGTCTAAA	2242
2297	AGACUGUC A UGCUAAUG	541	CATTAGCA GGCTAGCTACAACGA GACAGTCT	2243
2299	ACUGUCAU G CUAUUGGU	542	ACCAATTAG GGCTAGCTACAACGA ATGACAGT	2244
2303	UCAUGCUA A UGGUGUCC	543	GGACACCA GGCTAGCTACAACGA TAGCATGA	2245
2306	UGCUAAUG G UGUCCCCG	544	CGGGGACA GGCTAGCTACAACGA CATTAGCA	2246
2308	CUAUUGGU G UCCCCGAG	545	CTCGGGGA GGCTAGCTACAACGA ACCATTAG	2247
2316	GUCCCCGA G CCUCAGAU	546	ATCTGAGG GGCTAGCTACAACGA TCGGGGAC	2248
2323	AGCCUCAG A UCACUUGG	547	CCAAGTGA GGCTAGCTACAACGA CTGAGGCT	2249
2326	CUCAGAUC A CUUGGUUU	548	AAACCAAG GGCTAGCTACAACGA GATCTGAG	2250
2331	AUCACUUG G UUUAAAAA	549	TTTTTAAA GGCTAGCTACAACGA CAAGTGAT	2251
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2342	AAAAAAACA A CCACAAAA	551	TTTTGTGG GGCTAGCTACAACGA TGTTTTTA	2253
2345	AAACAAACC A CAAAUC	552	GTATTTTG GGCTAGCTACAACGA GGTTGTTT	2254
2350	ACCACAAA A UACAAACAA	553	TTGTTGTA GGCTAGCTACAACGA TTTGTTGT	2255
2352	CACAAAUU A CAACAAGA	554	TCTTGTG GGCTAGCTACAACGA ATTTTGTG	2256
2355	AAAAUACA A CAAGAGCC	555	GGCTCTTG GGCTAGCTACAACGA TGTATTTT	2257
2361	CAACAAGA G CCUGGAAU	556	ATTCCAGG GGCTAGCTACAACGA TCTTGTG	2258
2368	AGCCUGGA A UUAAUAAA	557	TAAATAAA GGCTAGCTACAACGA TCCAGGCT	2259
2371	CUGGAADU A UUUUAGGA	558	TCCTAAAA GGCTAGCTACAACGA AATTCCAG	2260
2379	AUUUUAGG A CCAGGAAG	559	CITCTCTGG GGCTAGCTACAACGA CCTAAAAT	2261
2387	ACCAGGAA G CAGCACGC	560	GCGTGCTG GGCTAGCTACAACGA TTCCCTGGT	2262
2390	AGGAAGCA G CACGCUGU	561	ACAGCGTG GGCTAGCTACAACGA TGCTTCCT	2263
2392	GAAGCAGC A CGCUGUUU	562	AAACAGCG GGCTAGCTACAACGA GCTGCTTC	2264
2394	AGCAGCAC G CUGUUUUA	563	ATAAACAG GGCTAGCTACAACGA GTGCTGCT	2265
2397	AGCACCGU G UUUAUJUGA	564	TCAATAAA GGCTAGCTACAACGA AGCGTGCT	2266
2401	CCCGUGUU A UUGAAAGA	565	TCTTTCAA GGCTAGCTACAACGA AAACAGCG	2267
2410	UUGAAAGA G UCACAGAA	566	TTCTGTGA GGCTAGCTACAACGA TCTTTCAA	2268
2413	AAAGAGUC A CAGAAGAG	567	CTCTTCTG GGCTAGCTACAACGA GACTCTTT	2269
2423	AGAAGAGG A UGAAGGUG	568	CACCTTCA GGCTAGCTACAACGA CCTCTTCT	2270

2429	GGAUGAAG G UGUCUAUC	569	GATAGACA GGCTAGCTACAACGA CTTCATCC	2271
2431	AUGAAGGU G UCUAUCAC	570	GTGATAGA GGCTAGCTACAACGA ACCTTCAT	2272
2435	AGGUGUCU A UCACUGCA	571	TGCAGTGA GGCTAGCTACAACGA AGACACCT	2273
2438	UGUCUAUC A CUGCAAAG	572	CTTTGCAG GGCTAGCTACAACGA GATAGACA	2274
2441	CUAUCACU G CAAAGCCA	573	TGGCTTIG GGCTAGCTACAACGA AGTGATAG	2275
2446	ACUGCAAA G CCACCAAC	574	GTTGGTGG GGCTAGCTACAACGA TTTGCAGT	2276
2449	GCAAAGCC A CCAACCAG	575	CTGGTTGG GGCTAGCTACAACGA GGCTTTGC	2277
2453	AGCCACCA A CCAGAAGG	576	CCTTCTGG GGCTAGCTACAACGA TGGTGGCT	2278
2462	CCAGAAGG G CUCUGUGG	577	CCACAGAG GGCTAGCTACAACGA CCTTCTGG	2279
2467	AGGGCUCU G UGGAAAGU	578	ACTTTCCA GGCTAGCTACAACGA AGAGCCCT	2280
2474	UGUGGAAA G UUCAGCAU	579	ATGCTGAA GGCTAGCTACAACGA TTTCCACA	2281
2479	AAAGUUC A CAUACCUC	580	GAGGTATG GGCTAGCTACAACGA TGAACTTT	2282
2481	AGUUCAGC A UACCUCAC	581	GTGAGGTA GGCTAGCTACAACGA GCTGAACT	2283
2483	UUCAGCAU A CCUCACUG	582	CAGTGAGG GGCTAGCTACAACGA ATGCTGAA	2284
2488	CAUACCUC A CUGUUCAA	583	TTGACACG GGCTAGCTACAACGA GAGGTATG	2285
2491	ACCUACACU G UUCAAGGA	584	TCCCTGAA GGCTAGCTACAACGA AGTGAGGT	2286
2500	UUCAAGGA A CCUCGGAC	585	GTCCGAGG GGCTAGCTACAACGA TCCCTGAA	2287
2507	AACCUCCG A CAAGUCUA	586	TAGACTTG GGCTAGCTACAACGA CCGAGGTT	2288
2511	UCGGACAA G UCUAAUCU	587	AGATTAGA GGCTAGCTACAACGA TTGTCCGA	2289
2516	CAAGUCUA A UCUGGAGC	588	GCTCCAGA GGCTAGCTACAACGA TAGACTTG	2290
2523	AAUCUGGA G CUGAUCAC	589	GTGATCAT GGCTAGCTACAACGA TCCAGATT	2291
2527	UGGAGCUG A UCACUCUA	590	TAGAGTGA GGCTAGCTACAACGA CAGCTCCA	2292
2530	AGCUGAUC A CUCUAAACA	591	TGTTAGAG GGCTAGCTACAACGA GATCAGCT	2293
2536	UCACUCUA A CAUGCACC	592	GGTGCATG GGCTAGCTACAACGA TAGAGTGA	2294
2538	ACUCUAAAC A UGCACCUG	593	CAGGTGCA GGCTAGCTACAACGA GTTAGAGT	2295
2540	UCUAACAU G CACCUGUG	594	CACAGGTG GGCTAGCTACAACGA ATGTTAGA	2296
2542	UAACAUAGC A CCUGUGUG	595	CACACAGG GGCTAGCTACAACGA GCATGTTA	2297
2546	AUGCACCU G UGUGGCUG	596	CAGCCACA GGCTAGCTACAACGA AGGTGCAT	2298
2548	GCACCUUG G UGGCUGCG	597	CGCAGCCA GGCTAGCTACAACGA ACAGGTGC	2299
2551	CCUGUGUG G CUGCGACU	598	AGTCGCAG GGCTAGCTACAACGA CACACAGG	2300
2554	GUGUGGCC G CGACUCUC	599	GAGAGTCG GGCTAGCTACAACGA AGCCACAC	2301
2557	UGGCUGCG A CUCUCUUC	600	GAAGAGAG GGCTAGCTACAACGA CGCAGCCA	2302
2568	CUCUUCUG G CUCCUAUJ	601	AATAGGAG GGCTAGCTACAACGA CAGAAGAG	2303
2574	UGGCUCU A UUAACCUU	602	AGGGTTAA GGCTAGCTACAACGA AGGAGCCA	2304
2578	UCCUAAUA A CCCUCUU	603	AAGGAGGG GGCTAGCTACAACGA TAATAGGA	2305
2587	CCCUCUUA A UCCGAAAA	604	TTTTCGGA GGCTAGCTACAACGA AAGGAGGG	2306
2596	UCCGAAAA A UGAAAAGG	605	CCTTTTCA GGCTAGCTACAACGA TTTTCGGA	2307
2604	AUGAAAAG G UCUUUCUUC	606	GAAGAAGA GGCTAGCTACAACGA CTTTTCAT	2308
2617	CUUCUGAA A UAAAAGACU	607	AGTCCTTA GGCTAGCTACAACGA TTCAGAAG	2309
2623	AAAUAAG A CUGACUAC	608	GTAGTCAG GGCTAGCTACAACGA CTTTATTI	2310
2627	AAAGACUG A CUACCUAJ	609	ATAGGTAG GGCTAGCTACAACGA CAGTCITI	2311
2630	GACUGACU A CCUAUCAA	610	TTGATAGG GGCTAGCTACAACGA AGTCAGTC	2312
2634	GACUACCU A UCAAAUUAJ	611	ATAATTGA GGCTAGCTACAACGA AGGTAGTC	2313
2638	ACCUAUCA A UUAAAUG	612	CATTATAA GGCTAGCTACAACGA TGATAGGT	2314
2641	UAUCAAUU A UAAUAGGAC	613	GTCCATTA GGCTAGCTACAACGA AAITTGATA	2315
2644	CAAUUAAA A UGGACCCA	614	TGGGTCCA GGCTAGCTACAACGA TATAATTG	2316
2648	UAAUAAUGG A CCCAGAUG	615	CATCTGGG GGCTAGCTACAACGA CCATTATA	2317
2654	GGACCCAG A UGAAGUUC	616	GAACCTCA GGCTAGCTACAACGA CTGGGTCC	2318
2659	CAGAUGAA G UUCCUUUG	617	CAAAGGAA GGCTAGCTACAACGA TTCACTG	2319
2669	UCCUUUUGG A UGAGCAGU	618	ACTGCTCA GGCTAGCTACAACGA CCAAAGGA	2320
2673	UUGGAUGA G CAGUGUGA	619	TCACACTG GGCTAGCTACAACGA TCATCCAA	2321
2676	GAUGAGCA G UGUGAGCG	620	CGCTCACAA GGCTAGCTACAACGA TGCTCATC	2322

2678	UGAGCAGU G UGAGCGGC	621	GCCGCTCA GGCTAGCTACAACGA ACTGCTCA	2323
2682	CAGUGUGA G CGGCCUCCC	622	GGGAGCCG GGCTAGCTACAACGA TCACACTG	2324
2685	UGUGAGCG G CUCCCCUUA	623	TAAGGGAG GGCTAGCTACAACGA CGCTCACA	2325
2693	GUCCCCUU A UGAUGCCA	624	TGGCATCA GGCTAGCTACAACGA AAGGGAGC	2326
2696	CCCUUAUG A UGCCAGCA	625	TGCTGGCA GGCTAGCTACAACGA CATAAGGG	2327
2698	CUUAUGAU G CCAGCAAG	626	CTTGCTGG GGCTAGCTACAACGA ATCATAAG	2328
2702	UGAUGCCA G CAAGUGGG	627	CCCACTTG GGCTAGCTACAACGA TGGCATCA	2329
2706	GCCAGCAA G UGGGGAGU	628	AACTCCCA GGCTAGCTACAACGA TTGCTGGC	2330
2712	AAGUGGGG G UUUGCCCG	629	CGGGCAAA GGCTAGCTACAACGA TCCCCATT	2331
2716	GGGAGUUU G CCCGGGAG	630	CTCCCGGG GGCTAGCTACAACGA AAACTCCC	2332
2727	CGGGAGAG A CUUAAAUC	631	AGTTTAAG GGCTAGCTACAACGA CTCTCCCG	2333
2733	AGACUAAA A CUGGGCAA	632	TTGCCCCAG GGCTAGCTACAACGA TTAAGTCT	2334
2738	UAAACUGG G CAAAUAC	633	GTGATTTG GGCTAGCTACAACGA CCAGTTTA	2335
2742	CUGGGCAA A UCACUUGG	634	CCAAGTGA GGCTAGCTACAACGA TTGCCCCAG	2336
2745	GGCAAAUC A CUUUGGAAG	635	CTTCCAAAG GGCTAGCTACAACGA GATTTGCC	2337
2758	GAAGAGGG G CUUUUGGA	636	TCCAAAAG GGCTAGCTACAACGA CCCTCTTC	2338
2770	UJGGAAAA G UGGUUCAA	637	TTGAACCA GGCTAGCTACAACGA TTTTCCAA	2339
2773	AAAAAGUG G UUCAAGCA	638	TGCTTGAA GGCTAGCTACAACGA CACTTTTC	2340
2779	UGGUUCAA G CAUCAGCA	639	TGCTGATG GGCTAGCTACAACGA TTGAACCA	2341
2781	GUUCAAGC A UCAGCAAU	640	AATGCTGA GGCTAGCTACAACGA GCTTGAAC	2342
2785	AAGCAUCA G CAUJUUGGC	641	GCCAAATG GGCTAGCTACAACGA TGATGCTT	2343
2787	GCAUCAGC A UUUGGCAU	642	ATGCCAAA GGCTAGCTACAACGA GCTGATGC	2344
2792	AGCAUUUG G CAUUAAGA	643	TCTTAATG GGCTAGCTACAACGA CAAATGCT	2345
2794	CAUUUGGC A UUAAGAAA	644	TTTCTTAA GGCTAGCTACAACGA GCCAAATG	2346
2802	AUUAAGAA A UCACCUAC	645	GTAGGTGA GGCTAGCTACAACGA TTCTTAAT	2347
2805	AAGAAAUC A CCUACGUG	646	CACGTAGG GGCTAGCTACAACGA GATTTCTT	2348
2809	AAUCACCU A CGUGCCGG	647	CCGGCACG GGCTAGCTACAACGA AGGTGATT	2349
2811	UCACCUAC G UGCCGGAC	648	GTCCGGCA GGCTAGCTACAACGA GTAGGTGA	2350
2813	ACCUACGU G CGGGACUG	649	CAGTCGG GGCTAGCTACAACGA ACGTAGGT	2351
2818	CGUGCCGG A CUGUGGGU	650	AGCCACAG GGCTAGCTACAACGA CGGGCACG	2352
2821	GCCGGACU G UGGCUGUG	651	CACAGCCA GGCTAGCTACAACGA AGTCCGGC	2353
2824	GGACUGUG G CUGUGAAA	652	TTTCACAG GGCTAGCTACAACGA CACAGTCC	2354
2827	CUGUGGCC G UGAAAAUG	653	CATTTTCA GGCTAGCTACAACGA AGCCACAG	2355
2833	CUGUGAAA A UGCUGAAA	654	TTTCAGCA GGCTAGCTACAACGA TTTCACAG	2356
2835	GUGAAAAU G CUGAAAGA	655	TCTTTTCAG GGCTAGCTACAACGA ATTTTCAC	2357
2848	AAGAGGGG G CCACGGCC	656	GGCCGTGG GGCTAGCTACAACGA CCCCTCTT	2358
2851	AGGGGGCC A CGGCCAGC	657	GCTGGCCG GGCTAGCTACAACGA GGCCCCCT	2359
2854	GGGCCACG G CCAGCGAG	658	CTCGCTGG GGCTAGCTACAACGA CGTGGCCC	2360
2858	CACGGCCA G CGAGUACA	659	TGTACTCG GGCTAGCTACAACGA TGGCCGTG	2361
2862	GCCAGCGA G UACAAAGC	660	GCTTTGTA GGCTAGCTACAACGA TCGCTGGC	2362
2864	CAGCGAGU A CAAAGCUC	661	GAGCTTTG GGCTAGCTACAACGA ACTGCTG	2363
2869	AGUACAAA G CUCUGAUG	662	CATCAGAG GGCTAGCTACAACGA TTTGTACT	2364
2875	AAGCUCUG A UGACUGAG	663	CTCAGTPCA GGCTAGCTACAACGA CAGAGCTT	2365
2878	CUCUGAUG A CUGAGCUA	664	TAGCTCA GGCTAGCTACAACGA CATCAGAG	2366
2883	AUGACUGA G CUAAAAAU	665	ATTTTTAG GGCTAGCTACAACGA TCAGTCAT	2367
2890	AGCUAAAA A UCUUGACC	666	GGTCAAGA GGCTAGCTACAACGA TTTTAGCT	2368
2896	AAAUCUUG A CCCACAUU	667	AATGTGGG GGCTAGCTACAACGA CAAGATTT	2369
2900	CUUGACCC A CAUJUGGC	668	GGCCAATG GGCTAGCTACAACGA GGGCAAG	2370
2902	UGACCCAC A UGGGCCAC	669	GTGGCCAA GGCTAGCTACAACGA GTGGGTCA	2371
2906	CCACAUUG G CCACCAUC	670	GATGGTGG GGCTAGCTACAACGA CAATGTGG	2372
2909	CAUJUGGC A CCAUCUGA	671	TCAGATGG GGCTAGCTACAACGA GGCCAATG	2373
2912	UGGCCACC A UCUGAACG	672	CGTTCAAGA GGCTAGCTACAACGA GGTGGCCA	2374

2918	CCAUCUGA A CGUGGUUA	673	TAACCACG GGCTAGCTACAACGA TCAGATGG	2375
2920	AUCUGAAC G UGGUUUAC	674	GTTAACCA GGCTAGCTACAACGA TTTCAGAT	2376
2923	UGAACGUG G UUAACCUG	675	CAGTTAA GGCTAGCTACAACGA CACGTTCA	2377
2927	CGUGGUUA A CCUGCUGG	676	CCAGCAGG GGCTAGCTACAACGA TAACCACG	2378
2931	GUUAACCU G CUGGGAGC	677	GCTCCCAG GGCTAGCTACAACGA AGGTTAAC	2379
2938	UGCUGGGG G CCUGCACC	678	GGTGCAGG GGCTAGCTACAACGA TCCCAGCA	2380
2942	GGGAGGCC G CACCAAGC	679	GCTGGTG GGCTAGCTACAACGA AGGCTCCC	2381
2944	GAGCCUGC A CCAAGCAA	680	TTGCTTGG GGCTAGCTACAACGA GCAGGCTC	2382
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2968	CUCUGAUG G UGAUUGUU	684	AACAATCA GGCTAGCTACAACGA CATCAGAG	2386
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2979	AUUGUUGA A UAUCUGAA	687	TTGCAGTA GGCTAGCTACAACGA TCAACAAT	2389
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2996	AUAUGGAA A UCUCUCCA	692	TGGAGAGA GGCTAGCTACAACGA TTCCATAT	2394
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3008	CUCCAACU A CCUCAAGA	694	TCTTGAGG GGCTAGCTACAACGA AGTTGGAG	2396
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3047	CAACAAGG A UGCAGCAC	701	GTGCTGCA GGCTAGCTACAACGA CCTTGTG	2403
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3127	UAGAUAGC G UCACCAGC	718	GCTGGTGA GGCTAGCTACAACGA GCTATCTA	2420
3130	AAAGCGUC A CCAGCAGC	719	GCTGCTGG GGCTAGCTACAACGA GACGCTAT	2421
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3137	CACCAAGCA G CGAAAGCU	721	AGCTTTCG GGCTAGCTACAACGA TGCTGGTG	2423
3143	CAGCGAAA G CUDUGCGA	722	TCGCAAAAG GGCTAGCTACAACGA TTTCCGCTG	2424
3148	AAAGCUUU G CGAGCUCC	723	GGACCTCG GGCTAGCTACAACGA AAAGCTTT	2425
3152	CUUUGCGA G CUCCGGCU	724	AGCCGGAG GGCTAGCTACAACGA TCGCAAAAG	2426

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3176	AGAUAAAA G UCUGAGUG	727	CACTCAGA GGCTAGCTACAACGA TTTTATCT	2429
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3275	GGCCAGAG G CAUGGAGU	745	ACTCCATG GGCTAGCTACAACGA CTCTGGCC	2447
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3322	ACCUUGCA G CGAGAAC	755	GTTTCTCG GGCTAGCTACAACGA TGCCAGGT	2457
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3347	AUCUGAGA A CAACGUGG	759	CCACGTTG GGCTAGCTACAACGA TCTCAGAT	2461
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3361	UGGUGAAG A UUUGUGAU	763	ATCACAAA GGCTAGCTACAACGA CTTCACCA	2465
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3368	GAUUUGUG A UUUUGGCC	765	GGCCAAAA GGCTAGCTACAACGA CACAAATC	2467
3374	UGAUUUUG G CCUUGGCC	766	GGGCAAGG GGCTAGCTACAACGA CAAAATCA	2468
3379	UUGGCCUU G CCCGGGAU	767	ATCCCGGG GGCTAGCTACAACGA AAGGCCAA	2469
3386	UGCCCGGG A UAUUUAUA	768	TATAAATA GGCTAGCTACAACGA CCCGGCA	2470
3388	CCCGGGAU A UUUUAUAG	769	CTTATAAA GGCTAGCTACAACGA ATCCCGGG	2471
3392	GGAAUAAA A UAAGAAC	770	GGTTCTTA GGCTAGCTACAACGA AAATATCC	2472
3398	UUAUAAA A CCCCCGAU	771	AATCGGGG GGCTAGCTACAACGA TCTTATAA	2473
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3407	CCCCGAU A UGUGAGAA	773	TTCTCACA GGCTAGCTACAACGA AATCGGGG	2475
3409	CCGAUUUA G UGAGAAAA	774	TTTTCTCA GGCTAGCTACAACGA ATAATCGG	2476
3422	AAAAGGAG A UACUUGAC	775	GTCGAGTA GGCTAGCTACAACGA CTCCCTTT	2477
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3441	CCUCUGAA A UGGAUUGC	778	GCCATCCA GGCTAGCTACAACGA TTCAGAGG	2480
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3456	GCUCCCGA A UCUAUCUU	781	AAGATAGA GGCTAGCTACAACGA TCGGGAGC	2483
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3481	UCUACAGC A CCAAGAGC	787	GCTCTTGG GGCTAGCTACAACGA GCTGTAGA	2489
3488	CACCAAGA G CGACGUGU	788	ACACGTCG GGCTAGCTACAACGA TCTTGGTG	2490
3491	CAAGAGCG A CGUGUGGU	789	ACCACACG GGCTAGCTACAACGA CGCTCTG	2491
3493	AGAGCGAC G UGUGGUCU	790	AGACCACCA GGCTAGCTACAACGA GTCGCTCT	2492
3495	AGCGACGU G UGGUCUUA	791	TAAGACCA GGCTAGCTACAACGA ACGTCGCT	2493
3498	GACGUGUG G UCUUACGG	792	CCGTAAGA GGCTAGCTACAACGA CACACGTC	2494
3503	GUGGUUUU A CGGAGUAU	793	ATACTCCG GGCTAGCTACAACGA AAGACCAC	2495
3508	CUUACCGA G UAUUGCUG	794	CAGCAATA GGCTAGCTACAACGA TCCGTAAG	2496
3510	UACGGAGU A UUGGUGUG	795	CACAGCAA GGCTAGCTACAACGA ACTCCGTA	2497
3513	GGAGUAUU G CUGUGGGG	796	TCCCACAG GGCTAGCTACAACGA AATACTCC	2498
3516	GUAUUGCU G UGGGAAAU	797	ATTTCCCA GGCTAGCTACAACGA AGCAATAC	2499
3523	UGUGGGAA' A UCUUUCUCC	798	GGAGAAGA GGCTAGCTACAACGA TTCCCCACA	2500
3536	CUCCUUAG G UGGGUCUC	799	GAGACCCA GGCTAGCTACAACGA CTAAGGAG	2501
3540	UUAGGUGG G UCUCCAUU	800	TATGGAGA GGCTAGCTACAACGA CCACCTAA	2502
3546	GGGUCUCC A UACCCAGG	801	CCTGGGTA GGCTAGCTACAACGA GGAGACCC	2503
3548	GUCUCCAU A CCCAGGAG	802	CTCCTGGG GGCTAGCTACAACGA ATGGAGAC	2504
3556	ACCCAGGA G UACAAAUG	803	CATTGTGA GGCTAGCTACAACGA TCCCTGGT	2505
3558	CCAGGAGU A CAAAUGGA	804	TCCATTTG GGCTAGCTACAACGA ACTCCCTGG	2506
3562	GAGUACAA A UGGAUGAG	805	CTCATCCA GGCTAGCTACAACGA TTGTACTC	2507
3566	ACAAAUGG A UGAGGACU	806	AGTCCTCA GGCTAGCTACAACGA CCATTGTT	2508
3572	GGAGUGGG A CUUUUGCA	807	TGCAAAAG GGCTAGCTACAACGA CCTCATCC	2509
3578	GGACUUUU G CAGUCGCC	808	GGCGACTG GGCTAGCTACAACGA AAAAGTCC	2510
3581	CUUUGCA G UCGCCUGA	809	TCAGGGCA GGCTAGCTACAACGA TGCAAAAG	2511
3584	UUGCAGUC G CCUGAGGG	810	CCCTCAGG GGCTAGCTACAACGA GACTGCAA	2512
3596	GAGGAAG G CAUGAGGA	811	TCCTCATG GGCTAGCTACAACGA CTTCCCTC	2513
3598	GGGAAGGC A UGAGGAUG	812	CATCTICA GGCTAGCTACAACGA GCCTTCCC	2514
3604	GCAUGAGG A UGAGAGCU	813	AGCTCTCA GGCTAGCTACAACGA CCTCATGC	2515
3610	GGAGUGAGA G CUCCUGAG	814	CTCAGGAG GGCTAGCTACAACGA TCTCATCC	2516
3618	GCUCUUGA G UACUCUAC	815	GTAGAGTA GGCTAGCTACAACGA TCAGGAGC	2517
3620	UCCUGAGU A CUCUACUC	816	GAGTAGAG GGCTAGCTACAACGA ACTCAGGA	2518
3625	AGUACUCU A CUCCUGAA	817	TTCAGGAG GGCTAGCTACAACGA AGAGTACT	2519
3634	CUCCUGAA A UCUCUACG	818	CTGATAGA GGCTAGCTACAACGA TTCAGGAG	2520
3638	UGAAAUCU A UCAGAUCA	819	TGATCTGA GGCTAGCTACAACGA AGATTTCA	2521
3643	UCUUAUCAG A UCAUGCUG	820	CAGCATGA GGCTAGCTACAACGA CTGATAGA	2522
3646	AUCAGAUCA UGGUGGAC	821	GTCCAGCA GGCTAGCTACAACGA GATCTGAT	2523
3648	CAGAUCAU G CUGGACUG	822	CAGTCCAG GGCTAGCTACAACGA ATGATCTG	2524
3653	CAUGCUGG A CUGGUGGC	823	GCCAGCG AGCTAGCTACAACGA CCAGCATG	2525
3656	GCUGGACU G CUGGCACA	824	TGTGCCAG GGCTAGCTACAACGA AGTCCAGC	2526
3660	GACUGGUG G CACAGAGA	825	TCTCTGTG GGCTAGCTACAACGA CAGCAGTC	2527
3662	CUGCUGGC A CAGAGACC	826	GGTCTCTG GGCTAGCTACAACGA GCCAGCAG	2528
3668	GCACAGAG A CCCAAAAG	827	CTTTTGGG GGCTAGCTACAACGA CTCTGTGC	2529
3681	AAAGAAAAG G CCAAGAUU	828	AACTCTGG GGCTAGCTACAACGA CTTTCCTT	2530

3687	AGGCCAAG A UUUGCAGA	829	TCTGCAAA GGCTAGCTACAACGA CTTGGCCT	2531
3691	CAAGAUUU G CAGAACUU	830	AAGTTCTG GGCTAGCTACAACGA AAATCTTG	2532
3696	UUUGCAGA A CUUGUGGA	831	TCCACAAG GGCTAGCTACAACGA TCTGAAA	2533
3700	CAGAACUU G UGGAAAAA	832	TTTTTCCA GGCTAGCTACAACGA AAGTTCTG	2534
3708	GUGGAAAA A CUAGGUGA	833	TCACCTAG GGCTAGCTACAACGA TTTTCCAC	2535
3713	AAAACUAG G UGAUUUGC	834	GCAAATCA GGCTAGCTACAACGA CTAGTTT	2536
3716	ACUAGGUG A UUUGCUUC	835	GAAGAAA GGCTAGCTACAACGA CACCTAGT	2537
3720	GGUGAUUU G CUUCAAGC	836	GCTTGAAG GGCTAGCTACAACGA AAATCACC	2538
3727	UGCUUCAA G CAAAGUUA	837	TACATTG GGCTAGCTACAACGA TTGAAGCA	2539
3731	UCAAGCAA A UGUACAAAC	838	GTTGTACA GGCTAGCTACAACGA TTGCTTGA	2540
3733	AAGCAAAU G UACAAACAG	839	CTGTTGTA GGCTAGCTACAACGA ATTTGTT	2541
3735	GCAAAUGU A CAACAGGA	840	TCCTGTTG GGCTAGCTACAACGA ACATTTGC	2542
3738	AAUGUACA A CAGGAUGG	841	CCATCCTG GGCTAGCTACAACGA TGTACATT	2543
3743	ACAACAGG A UGGUAAAAG	842	CTTTACCA GGCTAGCTACAACGA CCTGTTGT	2544
3746	ACAGGAUO G UAAAGACU	843	AGTCTTTA GGCTAGCTACAACGA CATCCGT	2545
3752	UGGUAAAAG A CUACAUCC	844	GGATGTAG GGCTAGCTACAACGA CTTTACCA	2546
3755	UAAAGACU A CAUCCCAA	845	TTGGGATG GGCTAGCTACAACGA AGTCTTTA	2547
3757	AAGACUAC A UCCCCAAC	846	GATTGGGA GGCTAGCTACAACGA GTAGTCTT	2548
3763	ACAUCCCA A UCAAUGCC	847	GGCAITGA GGCTAGCTACAACGA TGGGATGT	2549
3767	CCCAAUCA A UGCCAUAC	848	GTATGGCA GGCTAGCTACAACGA TGATTGGG	2550
3769	CAAUCAAU G CCAUACUG	849	CAGTATGG GGCTAGCTACAACGA ATTGATTG	2551
3772	UCAAUGCC A UACUGACA	850	TGTCAAGT GGCTAGCTACAACGA GGCATTGA	2552
3774	AAUGCCAU A CUGACAGG	851	CCTGTCAG GGCTAGCTACAACGA ATGGCATT	2553
3778	CCAUACUG A CAGGAAAU	852	ATTTCTTG GGCTAGCTACAACGA CAGTATGG	2554
3785	GACAGGAA A UAGUGGGU	853	ACCCACTA GGCTAGCTACAACGA TTCCCTGTC	2555
3788	AGGAAAUU G UGGGUUUA	854	TAAACCCA GGCTAGCTACAACGA TATTTCTT	2556
3792	AAUAGUGG G UUUACAUU	855	TATGTAAA GGCTAGCTACAACGA CCACTATT	2557
3796	UGGGGUUU A CAUACUCA	856	TGAGTATG GGCTAGCTACAACGA AAACCCAC	2558
3798	GGGUUUUAC A UACUCAAC	857	GTTGAGTA GGCTAGCTACAACGA GTAAACCC	2559
3800	GUUUACAU A CUCAACUC	858	GAGTTGAG GGCTAGCTACAACGA ATGTAAAC	2560
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3811	CAACUCCU G CCUUCUCU	860	AGAGAAGG GGCTAGCTACAACGA AGGAGTTG	2562
3824	CUCUGAGG A CUUCUUCA	861	TGAAGAAG GGCTAGCTACAACGA CCTCAGAG	2563
3839	CAAGGAAA G UAUUUCAG	862	CTGAAATA GGCTAGCTACAACGA TTTCCTTG	2564
3841	AGGAAAGU A UUUCAGCU	863	AGCTGAAA GGCTAGCTACAACGA ACTTTCTT	2565
3847	GUAUUUCU G CUCCGAAG	864	CTTCGGAG GGCTAGCTACAACGA TGAAATAC	2566
3855	GCUCCGAA G UUUAAAUC	865	GAATTAAA GGCTAGCTACAACGA TTCCGGAGC	2567
3860	GAAGUUUA A UUCAGGAA	866	TTCCCTGAA GGCTAGCTACAACGA TAAACTTC	2568
3869	UUCAGGAA G CUCUGAUG	867	CATCAGAG GGCTAGCTACAACGA TTCCCTGAA	2569
3875	AAGCUCUG A UGAUGUCA	868	TGACATCA GGCTAGCTACAACGA CAGAGCTT	2570
3878	CUCUGAUG A UGUCAGAU	869	ATCTGACA GGCTAGCTACAACGA CATCAGAG	2571
3880	CUGAUGAU G UCAGAUU	870	ATATCTGA GGCTAGCTACAACGA ATCATCAG	2572
3885	GAUGUCAG A UAUUAAA	871	TTTACATA GGCTAGCTACAACGA CTGACATC	2573
3887	UGUCAGAU A UGUAAAUG	872	CATTTACA GGCTAGCTACAACGA ATCTGACA	2574
3889	UCAGAUAU G UAAAUGCU	873	AGCATTAA GGCTAGCTACAACGA ATATCTGA	2575
3893	AUAUGUAA A UGCUUUCU	874	TGAAAGCA GGCTAGCTACAACGA TTACATAT	2576
3895	AUGUAAAU G CUUUCAG	875	CTTGAAGA GGCTAGCTACAACGA ATTTACAT	2577
3903	GCUUUCAA G UUCAUGAG	876	CTCATGAA GGCTAGCTACAACGA TTGAAAGC	2578
3907	UCAAGUUC A UGAGCCUG	877	CAGGCTCA GGCTAGCTACAACGA GAACTTGA	2579
3911	GUUCAUGA G CCUGGAAA	878	TTTCCAGG GGCTAGCTACAACGA TCATGAAC	2580
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3939	UUUGAAGA A CUUUUACC	881	GGTAAAAG GGCTAGCTACAACGA TCTTCAAA	2583
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3950	UUUACCGA A UGCCACCU	883	AGGTGGCA GGCTAGCTACAACGA TCGGTAAA	2585
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3955	CGAAUGCC A CCUCCAUG	885	CATGGAGG GGCTAGCTACAACGA GGCAITCG	2587
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3971	GUUUGAUG A CUACCAAG	889	CCTGGTAG GGCTAGCTACAACGA CATCAAAC	2591
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3986	GGGCGACA G CAGCACUC	893	GAGTGCTG GGCTAGCTACAACGA TGTCGCCC	2595
3989	CGACAGCA G CACUCUGU	894	ACAGAGTG GGCTAGCTACAACGA TGCTGTCG	2596
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4030	UCACCUUG A CUGACAGC	903	GCTGTCAG GGCTAGCTACAACGA CCAGGTGA	2605
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4037	GACUGACA G CAAACCCA	905	TGGGTTTG GGCTAGCTACAACGA TGTCAGTC	2607
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4060	CGCUCAAG A UUGACUUG	909	CAAGTCAA GGCTAGCTACAACGA CTTGAGCG	2611
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4530	ACCCAAUG A CUUCCUG	1016	CAGGGAAAG GGCTAGCTACAACGA CATTGGGT	2718
4538	ACUUCUCC G CUCCAACC	1017	GGTTGGAG GGCTAGCTACAACGA AGGGAAGT	2719
4544	CUGCUCCA A CCCCCGCC	1018	GGCGGGGG GGCTAGCTACAACGA TGGAGCAG	2720
4550	CAACCCCC G CCACCUCA	1019	TGAGGTGG GGCTAGCTACAACGA GGGGGTTG	2721
4553	CCCCCGCC A CCUCAGGG	1020	CCCTGAGG GGCTAGCTACAACGA GGCGGGGG	2722
4561	ACCUCAGG G CACGCAGG	1021	CCTGGGTG GGCTAGCTACAACGA CCTGAGGT	2723
4563	CUCAGGGC A CGCAGGAC	1022	GTCCCTGG GGCTAGCTACAACGA GCCCTGAG	2724
4565	CAGGGCAC G TAGGACCA	1023	TGGTCCTG GGCTAGCTACAACGA GTGCCCTG	2725
4570	CACGCAGG A CCAGUUUG	1024	CAAACCTGG GGCTAGCTACAACGA CCTGCGTG	2726
4574	CAGGACCA G UUUGAUUG	1025	CAATCAAA GGCTAGCTACAACGA TGGTCCTG	2727
4579	CCAGUUUG A UUGAGGAG	1026	CTCCTCAA GGCTAGCTACAACGA CAAACTGG	2728
4587	AUUGAGGA G CUGCAUCU	1027	CAGTGCGAG GGCTAGCTACAACGA TCCCTCAAT	2729
4590	GAGGAGCU G CACUGAUC	1028	GATCACTG GGCTAGCTACAACGA AGCTCCTC	2730
4592	GGAGCUGC A CUGAUAC	1029	GTGATCAG GGCTAGCTACAACGA GCAGCTCC	2731
4596	CUGCACUG A UCACCCAA	1030	TTGGGTGA GGCTAGCTACAACGA CAGTGCAG	2732
4599	CACUGAUC A CCCAAUGC	1031	GCATTGGG GGCTAGCTACAACGA GATCAGTG	2733
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4606	CACCCAAU G CAUCACGU	1033	ACGTGATG GGCTAGCTACAACGA ATTGGGTG	2735
4608	CCCAAUGC A UCACGUAC	1034	GTACGTGA GGCTAGCTACAACGA GCATTGGG	2736
4611	AAUGCAUC A CGUACCCC	1035	GGGGTACG GGCTAGCTACAACGA GATGCATT	2737
4613	UGCAUCAC G UACCCAC	1036	GTGGGGTA GGCTAGCTACAACGA GTGATGCA	2738

4615	CAUCACGU A CCCCACUG	1037	CAGTGGGG GGCTAGCTACAACGA ACGTGATG	2739
4620	CGUACCCC A CUUCCCCA	1038	TGGCCAG GGCTAGCTACAACGA GGGGTACG	2740
4625	CCCACUGG G CCAGCCCU	1039	AGGGCTGG GGCTAGCTACAACGA CCAGTGGG	2741
4629	CUUCCCCA G CCCUGCAG	1040	CTGCAGGG GGCTAGCTACAACGA TGGCCAG	2742
4634	CCAGCCCU G CAGCCCAA	1041	TTGGGCTG GGCTAGCTACAACGA AGGGCTGG	2743
4637	GCCCCUGCA G CCCAAAC	1042	GTTTGGGG GGCTAGCTACAACGA TGCAAGGC	2744
4644	AGCCCCAA A CCCAGGGC	1043	GCCCTGGG GGCTAGCTACAACGA TTTGGGCT	2745
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4658	GGCAACAA G CCCGUUAG	1046	CTAACGGG GGCTAGCTACAACGA TTGTTGCC	2748
4662	ACAAGCCC G UUAGCCCC	1047	GGGGCTAA GGCTAGCTACAACGA GGGCTGT	2749
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4698	UGAGCAAC A UCUCGGGA	1055	TCCCGAGA GGCTAGCTACAACGA GTTGCTCA	2757
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4715	GUCCUCUA G CAGGCCUA	1057	TAGGCCTG GGCTAGCTACAACGA TAGAGGAC	2759
4719	UCUAGCAG G CCUAAGAC	1058	GTCTTAGG GGCTAGCTACAACGA CTGCTAGA	2760
4726	GGCCUUAAG A CAUGUGAG	1059	CTCACATG GGCTAGCTACAACGA CTTAGGCC	2761
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4814	GACGCACC A UGUGGGCA	1071	TGCCCAACA GGCTAGCTACAACGA GGTGCCCTC	2773
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4847	UCAGCAAU G CCAUUUCA	1079	TGAAATGG GGCTAGCTACAACGA ATTGCTGA	2781
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4859	UUUCAGUG G CUUCCCCA	1082	CTGGGAAG GGCTAGCTACAACGA CACTGAAA	2784
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4891	AUUUGAGG G CCCAGCCA	1087	TGGCTGGG GGCTAGCTACAACGA CCTCAAAT	2789
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4917	GGACAGCG A UGAGGGGA	1093	TCCCCTCA GGCTAGCTACAACGA CGCTGTCC	2795
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4927	GAGGGGAC A UUUUCUGG	1095	CCAGAAAA GGCTAGCTACAACGA GTCCCCTC	2797
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5609	GAAGAAAA G CCCAUUUU	1239	AAAATGGG GGCTAGCTACAACGA TTTTCTTC	2941
5613	AAAAGCCC A UUUUCAAC	1240	GTTGAAAA GGCTAGCTACAACGA GGGCTTTT	2942
5620	CAUUUUCU A CUGCUUUG	1241	CAAAGCAG GGCTAGCTACAACGA TGAAAATG	2943
5623	UUUCAACU G CUUUGAAA	1242	TTTCAAAG GGCTAGCTACAACGA AGTTGAAA	2944
5631	GUUUGGAA A CIIUGCCUG	1243	CAGGCAAG GGCTAGCTACAACGA TTCAAAGC	2945
5635	UGAAACUU G CCUGGGGU	1244	ACCCCAAGG GGCTAGCTACAACGA AAGTTICA	2946

5642	UGCCUGGG G UCUGAGCA	1245	TGCTCAGA GGCTAGCTACAACGA CCCAGGCA	2947
5648	GGGUCUGA G CAUGAUGG	1246	CCATCATG GGCTAGCTACAACGA TCAGACCC	2948
5650	GUCUGAGC A UGAUGGGA	1247	TCCCCTCA GGCTAGCTACAACGA GCTCAGAC	2949
5653	UGAGCAUG A UGGGAAUA	1248	TATTCCCA GGCTAGCTACAACGA CATGCTCA	2950
5659	UGAUGGGA A UAGGGAGA	1249	TCTCCCTA GGCTAGCTACAACGA TCCCCTAT	2951
5667	AUAGGGAG A CAGGGUAG	1250	CTACCCCTG GGCTAGCTACAACGA CTCCCTAT	2952
5672	GAGACAGG G UAGGAAAG	1251	CTTTCCTA GGCTAGCTACAACGA CCTGTCTC	2953
5682	AGGAAAGG G CGCCUACU	1252	AGTAGGCG GGCTAGCTACAACGA CCTTTCCCT	2954
5684	GAAAGGGC G CCUACUCU	1253	AGAGTAGG GGCTAGCTACAACGA GCCCTTTC	2955
5688	GGGCGCCU A CUCUUCAG	1254	CTGAAGAG GGCTAGCTACAACGA AGGCGCCC	2956
5698	UCUUCAGG G UCUAAAGA	1255	TCTTTAGA GGCTAGCTACAACGA CCTGAAGA	2957
5706	GUCUAAAG A UCAAGUGG	1256	CCACTTG A GGCTAGCTACAACGA CTTTAGAC	2958
5711	AAGAUCAA G UGGGCCUU	1257	AAGGCCCA GGCTAGCTACAACGA TTGATCTT	2959
5715	UCAAGUGG G CCUUGGAU	1258	ATCCAAGG GGCTAGCTACAACGA CCACTTGA	2960
5722	GGCCUUGG A UCGCUAAG	1259	CTTACCGA GGCTAGCTACAACGA CCAAGGCC	2961
5725	CUUGGAUC G CUAAGCUG	1260	CAGCTTAG GGCTAGCTACAACGA GATCCAAG	2962
5730	AUCGCUAA G CUGGCUCU	1261	AGAGCCAG GGCTAGCTACAACGA TTAGCGAT	2963
5734	CUAAGCUG G CUCUGUUU	1262	AAACAGAG GGCTAGCTACAACGA CAGCTTAG	2964
5739	CUGGCUCU G UUUGAUGC	1263	GCATCAAA GGCTAGCTACAACGA AGAGCCAG	2965
5744	UCUGUUUG A UGCUAUUU	1264	AAATAGCA GGCTAGCTACAACGA CAAACAGA	2966
5746	UGUUUGAU G CUAUUUAU	1265	ATAAAATAG GGCTAGCTACAACGA ATCAAACA	2967
5749	UUGAUGCU A UUUUAGCA	1266	TGCATAAA GGCTAGCTACAACGA AGCATCAA	2968
5753	UGCUAUUU A UGCAAGUU	1267	AACTTGCA GGCTAGCTACAACGA AAATAGCA	2969
5755	CUAUUUAU G CAAGUUAG	1268	CTAACTTG GGCTAGCTACAACGA ATAAATAG	2970
5759	UUAUGCAA G UUAGGGUC	1269	GACCCCTAA GGCTAGCTACAACGA TTGCATAA	2971
5765	AAGUUAGG G UCUAUGUA	1270	TACATAGA GGCTAGCTACAACGA CCTTAACCTT	2972
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5782	UUUAGGAU G CGCCUACU	1275	AGTAGGCG GGCTAGCTACAACGA ATCCCTAA	2977
5784	UAGGAUGC G CCUACUCU	1276	AGAGTAGG GGCTAGCTACAACGA GCATCCTA	2978
5788	AUGCGCCU A CUCUUCAG	1277	CTGAAGAG GGCTAGCTACAACGA AGGCGCAT	2979
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5806	GUCUAAAG A UCAAGUGG	1279	CCACTTGA GGCTAGCTACAACGA CTTTAGAC	2981
5811	AAGAUCAA G UGGGCCUU	1280	AAGGCCCA GGCTAGCTACAACGA TTGATCTT	2982
5815	UCAAGUGG G CCUUGGAU	1281	ATCCAAGG GGCTAGCTACAACGA CCACTTGA	2983
5822	GGCCUUGG A UCGCUAAG	1282	CTTACCGA GGCTAGCTACAACGA CCAAGGCC	2984
5825	CUUGGAUC G CUAAGCUG	1283	CAGCTTAG GGCTAGCTACAACGA GATCCAAG	2985
5830	AUCGCUAA G CUGGCUCU	1284	AGAGCCAG GGCTAGCTACAACGA TTAGCGAT	2986
5834	CUAAGCUG G CUCUGUUU	1285	AAACAGAG GGCTAGCTACAACGA CAGCTTAG	2987
5839	CUGGCUCU G UUUGAUGC	1286	GCATCAAA GGCTAGCTACAACGA AGAGCCAG	2988
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5869	UAGGGUCU A UGUAUUUA	1294	TAAATACA GGCTAGCTACAACGA AGACCCCTA	2996
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5902	UGCAGCCA G UCAGAACG	1303	GCTTCTGA GGCTAGCTACAACGA TGGCTGCA	3005
5909	AGUCAGAA G CUGGAGAG	1304	CTCTCCAG GGCTAGCTACAACGA TTCTGACT	3006
5918	CUGGAGAG G CAACAGUG	1305	CACTGTTG GGCTAGCTACAACGA CTCTCCAG	3007
5921	GAGAGGCC A CAGUGGAU	1306	ATCCACTG GGCTAGCTACAACGA TGCCCTTC	3008
5924	AGGCAACA G UGGAUUGC	1307	GCAATCCA GGCTAGCTACAACGA TGTGCGCT	3009
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5931	AGUGGAUU G CUGCUUCU	1309	AGAAGCAG GGCTAGCTACAACGA AATCCACT	3011
5934	GGAUUGC G CUUCUUGG	1310	CCAAGAAG GGCTAGCTACAACGA AGCAATCC	3012
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5955	AAGAGUAU G CUUCCUUU	1313	AAAGGAAG GGCTAGCTACAACGA ATACTCTT	3015
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5987	ACUGUAGA A CCUGAGCU	1320	AGCTCAGG GGCTAGCTACAACGA TCTACAGT	3022
5993	GAACCUGA G CUCUAAGU	1321	ACTTAGAG GGCTAGCTACAACGA TCAGGTTTC	3023
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6003	UCUAAGUA A CCGAAGAA	1323	TTCTTCGG GGCTAGCTACAACGA TACTTACA	3025
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6069	AUGAAGAG A UGGGACCG	1338	CGGTCCCC GGCTAGCTACAACGA CTCTTCAT	3040
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6672	AAACAUAG A UUAACUGA	1460	TCAGTTAA GGCTAGCTACAACGA TCATGTTT	3162
6676	AUGAAUUA A CUGAUAAA	1461	ATTATCAAG GGCTAGCTACAACGA TAATTCAAT	3163
6680	AUUAACUG A UAAAUUUC	1462	GAATATTA GGCTAGCTACAACGA CAGTTAAT	3164
6683	AACUGAU A UAUUCCAA	1463	TTGGAAATA GGCTAGCTACAACGA TATCAGTT	3165
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6698	AAUCAUU G CCAUUUAU	1467	ATAAAATGG GGCTAGCTACAACGA AAATGATT	3169
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6705	UGCCAUU A UGACAAAA	1469	TTTTGTCA GGCTAGCTACAACGA AAATGGCA	3171
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6738	AAGAACGA G CACUUCU	1477	AGGAAGTG GGCTAGCTACAACGA TCGTTCTT	3179
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6753	CUUUCAGA G UUUCUGAG	1479	CTCAGAAA GGCTAGCTACAACGA TCTGAAAG	3181
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6767	GAGAUAAA G UACGUGGA	1482	TCCACGTA GGCTAGCTACAACGA ATTATCTC	3184
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6771	UAAUGUAC G UGGAACAG	1484	CTGTTCCA GGCTAGCTACAACGA GTACATTA	3186
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6779	UGGGAAACA G UCUGGGUG	1486	CACCCAGA GGCTAGCTACAACGA TGTTCCAC	3188
6785	CAGUCUGG G UGGAAUUG	1487	CCATTCCA GGCTAGCTACAACGA CCAGACTG	3189
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7563	AUCACCU A UGGAUAUU	1678	AATATCCA GGCTAGCTACAACGA AGGGTGAT	3380
7567	CCCUAUGG A UAUUGGU	1679	AGCCAATAA GGCTAGCTACAACGA CCATAGGG	3381
7569	CUAUGGAU A UUGGUAG	1680	CTAGCCAA GGCTAGCTACAACGA ATCCATAG	3382
7573	GGAUAUUG G CUAGUUU	1681	AAAATAG GGCTAGCTACAACGA CAATATCC	3383
7577	AUUGGCCA G UUUUGCCU	1682	AGGCAAAA GGCTAGCTACAACGA TAGCCAAT	3384
7582	CUAGUUUU G CCUUUUAU	1683	AATAAAGG GGCTAGCTACAACGA AAAACTAG	3385
7588	UUGCCUUU A UUAAGCAA	1684	TTGCTTAA GGCTAGCTACAACGA AAAGGCAA	3386
7593	UUUAUUA G CAAAUUCA	1685	TGAATTGG GGCTAGCTACAACGA TTAATAAA	3387
7597	UUAAGCAA A UUCAUUC	1686	GAATGAA GGCTAGCTACAACGA TTGCTTAA	3388
7601	GCAAAUUC A UUUCAGCC	1687	GGCTGAAA GGCTAGCTACAACGA GAATTTGC	3389
7607	UCAUUUCA G CCUGAANG	1688	CATTCAGG GGCTAGCTACAACGA TGAAATGA	3390
7613	CAGCCUGA A UGUCUGCC	1689	GGCAGACAA GGCTAGCTACAACGA TCAGGCTG	3391
7615	GCCUGAAU G UCUGCCUA	1690	TAGGCAGA GGCTAGCTACAACGA ATTCAAGGC	3392
7619	GAAUGUCU G CCUUAUUA	1691	TATATAGG GGCTAGCTACAACGA AGACATTC	3393
7623	GUCUGCCU A UAUUAUUC	1692	AGAATATAA GGCTAGCTACAACGA AGGCAGAC	3394
7625	CUGCCUAU A UAUUCUCU	1693	AGAGAATAA GGCTAGCTACAACGA ATAGGCAG	3395
7627	GCCUUAU A UUCUCUGC	1694	GCAGAGAAA GGCTAGCTACAACGA ATATAGGC	3396
7634	UAUUCUCU G CUCUUUGU	1695	ACAAAGAG GGCTAGCTACAACGA AGAGAATA	3397
7641	UGCUCUUU G UAUUCUCC	1696	GGAGAATAA GGCTAGCTACAACGA AAAGAGCA	3398
7643	CUCUUUGU A UUCUCCUU	1697	AAGGAGAAA GGCTAGCTACAACGA ACAAAAGAG	3399
7655	UCCUUUGA A CCCGUAAA	1698	TTAACCGGG GGCTAGCTACAACGA TCAAAGGA	3400
7659	UUGAACCC G UAAAAAAC	1699	TGTTTTAA GGCTAGCTACAACGA GGGTTCAA	3401
7665	CCGUAAAA A CAUCCUGU	1700	ACAGGATG GGCTAGCTACAACGA TTTAACGG	3402
7667	GUUAAAAC A UCCUGUGG	1701	CCACAGGA GGCTAGCTACAACGA GTTTTAAAC	3403
7672	AACAUCCU G UGGCACUC	1702	GAGTGCCCA GGCTAGCTACAACGA AGGATGTT	3404

Input Sequence = HSFLT. Cut Site = R/Y

Arm Length = 8. Core Sequence = GGCTAGCTACAACGA

HSFLT (Human flt mRNA for receptor-related tyrosine kinase.; Acc# X51602; 7680 bp)

Table VI: Human KDR DNAzyme and Substrate sequence

Pos	Substrate	Seq ID No	DNAzyme	Seq ID No
14	GUCCCCGG A CCCCGGGA	3405	TCCCGGGG GGCTAGCTACAACGA CCCGGGAC	4691
25	CCGGGAGA G CGGUUCAGU	3406	ACTGACCG GGCTAGCTACAACGA TCTCCCGG	4692
28	GGAGAGCG G UCAGUGUG	3407	CACACTGA GGCTAGCTACAACGA CGCTCTCC	4693
32	AGCCGUCA G UGUGUGGU	3408	ACCACACA GGCTAGCTACAACGA TGACCGCT	4694
34	CGGUAGU G UGUGGUUC	3409	CGACCACA GGCTAGCTACAACGA ACTGACCG	4695
36	GUCAAGU G UGGUCGCU	3410	AGCGACCA GGCTAGCTACAACGA ACAGTGAC	4696
39	AGUGUGUG G UCGCUGCG	3411	CGCAGCGA GGCTAGCTACAACGA CACACACT	4697
42	GUGUGGUC G CUCCGUUU	3412	AAACGCAG GGCTAGCTACAACGA GACCACAC	4698
45	UGGUUCGU G CGUUUUCU	3413	AGGAAACG GGCTAGCTACAACGA AGCGACCA	4699
47	GUCCUGC G UUUCUCU	3414	AGAGGAAA GGCTAGCTACAACGA GCAGCGAC	4700
56	UUUCCUCU G CCUGCGCC	3415	GGCGCAGG GGCTAGCTACAACGA AGAGGAAA	4701
60	CUCUCCU G CGCCGGGC	3416	GCCCGGGG GGCTAGCTACAACGA AGGCAGAG	4702
62	CUGCCUGC G CGGGCAU	3417	ATGCCCGG GGCTAGCTACAACGA GCAGGCAG	4703
67	UGGCCGG G CAUCACUU	3418	AAGTGATG GGCTAGCTACAACGA CGGGCGCA	4704
69	CGCCGGGC A UCACUUGC	3419	GCAAGTGA GGCTAGCTACAACGA GCCCGGGG	4705
72	CGGGCAUC A CUUGCGCG	3420	CGCGCAAG GGCTAGCTACAACGA GATGCCCG	4706
76	AUCACUU G CGCGCCGC	3421	GCGGCGCG GGCTAGCTACAACGA AAGTGATG	4707
78	UCACUUGC G CGCCGCAG	3422	CTGCGGCG GGCTAGCTACAACGA GCAAGTGA	4708
80	ACUUGCGC G CGCGAGAA	3423	TTCTGCGG GGCTAGCTACAACGA GCGCAAGT	4709
83	UGCCGCGC G CAGAAAAGU	3424	ACTTTCTG GGCTAGCTACAACGA GGCGCGCA	4710
90	CGCAGAAA G UCCGUCUG	3425	CAGACGGA GGCTAGCTACAACGA TTTCTGCG	4711
94	GAAAGUCC G UCUGGCAG	3426	CTGCCAGA GGCTAGCTACAACGA GGACTTTC	4712
99	UCCGUCUG G CGGCCUGG	3427	CCAGGCTG GGCTAGCTACAACGA CAGACGGA	4713
102	GUCUGGCA G CCUGGAAU	3428	TATCCAGG GGCTAGCTACAACGA TGCCAGAC	4714
108	CAGCCUGG A UAUCUCU	3429	AGAGGATA GGCTAGCTACAACGA CCAGGCTG	4715
110	GCCUGGAU A UCCUCUCC	3430	GGAGAGGA GGCTAGCTACAACGA ATCCAGGC	4716
120	CCUCUCCU A CGCCGCACC	3431	GGTGCGGG GGCTAGCTACAACGA AGGAGAGG	4717
124	UCCUACCG G CACCCGCA	3432	TGCGGGTG GGCTAGCTACAACGA CGGTAGGA	4718
126	CUACCGC A CCCGCAGA	3433	TCTGCGGG GGCTAGCTACAACGA GCGGGTAG	4719
130	CGGCACCC G CAGACGCC	3434	GGCGTCTG GGCTAGCTACAACGA GGGTGCGG	4720
134	ACCCGCAG A CGCCCCUG	3435	CAGGGGGG GGCTAGCTACAACGA CTGCGGGT	4721
136	CGCGAGAC G CCCCUGCA	3436	TGCAGGGG GGCTAGCTACAACGA GTCTGCGG	4722
142	ACGCCCU G CAGCCGCC	3437	GGCGGCTG GGCTAGCTACAACGA AGGGGCGT	4723
145	CCCCUGCA G CGCCGGGU	3438	ACCGGGCG GGCTAGCTACAACGA TGCAAGGG	4724
148	CUGCAGCC G CGGGUCGG	3439	CCGACCGG GGCTAGCTACAACGA GGCTGCAG	4725
152	AGCCGCG G UCGCGCGC	3440	GGCGCCGA GGCTAGCTACAACGA CGGGGGCT	4726
156	GCGGGUOG G CGCCCGGG	3441	CCCGGGCG GGCTAGCTACAACGA CGACCGGC	4727
158	CGGUCCGC G CGCGGGCU	3442	AGCCCGGG GGCTAGCTACAACGA GCCGACCG	4728
164	GCGCCCGG G CUCCCCUAG	3443	CTAGGGAG GGCTAGCTACAACGA CGGGCGC	4729
172	GCUCCCUA G CCCUGUGC	3444	GCACAGGG GGCTAGCTACAACGA TAGGGAGC	4730
177	CUTAGCCCCU G UGCGCUCA	3445	TGAGCGCA GGCTAGCTACAACGA AGGGCTAG	4731
179	ACCCCCUGU G CGCUAAC	3446	GTTGAGCG GGCTAGCTACAACGA ACAGGGCT	4732
181	CCCUGUGC G CUTAACUG	3447	CAGTTGAG GGCTAGCTACAACGA GCACAGGG	4733
186	UGCGCUCA A CUGUCCUG	3448	CAGGACAG GGCTAGCTACAACGA TGAGCGCA	4734
189	GCUCAACU G UCCUGCGC	3449	GCGCAGGA GGCTAGCTACAACGA AGTTGAGC	4735
194	ACUGUCCU G CGCUGCGG	3450	CGGCACCG GGCTAGCTACAACGA AGGACAGT	4736
196	UGUCCUGC G CGUGGGGG	3451	CCCCCGAG GGCTAGCTACAACGA GCAGGGACA	4737
199	CCUGCGCU G CGGGGUGC	3452	GCACCCCC GGCTAGCTACAACGA AGGGCAGG	4738
204	CGUGCGGG G UGCGCGCA	3453	TCGGCGCA GGCTAGCTACAACGA CCCGCAGC	4739

206	UGCGGGGU G CCGCGAGU	3454	ACTCGCGG GGCTAGCTACAACGA ACCCCGCA	4740
209	GGGGUGCC G CGAGUUCC	3455	GGAACTCG GGCTAGCTACAACGA GGCACCCC	4741
213	UGCCCGCA G UUCCACCU	3456	AGGTGGAA GGCTAGCTACAACGA TCGGGCA	4742
218	CGAGUUCC A CCUCGGCG	3457	CGCGGAGG GGCTAGCTACAACGA GGAACTCG	4743
224	CCACCUCC G CGCCUCCU	3458	AGGAGGCG GGCTAGCTACAACGA GGAGGTGG	4744
226	ACCUCCGC G CCUCUUC	3459	GAAGGAGG GGCTAGCTACAACGA GCGGAGGT	4745
240	UUCUCUAG A CAGGCGCU	3460	AGCGCCTG GGCTAGCTACAACGA CTAGAGAA	4746
244	CUAGACAG G CGCUGGGA	3461	TCCCAGCG GGCTAGCTACAACGA CTGTCTAG	4747
246	AGACAGGC G CUGGGAGA	3462	TCTCCCAG GGCTAGCTACAACGA GCCTGTCT	4748
259	GAGAAAGA A CCGGCCUCC	3463	GGAGCCGG GGCTAGCTACAACGA TCTTTCTC	4749
263	AAGAACCG G CUCCCCAG	3464	CTCAGGGAG GGCTAGCTACAACGA CGGTTCTT	4750
271	GUCCCCGA G UUCUGGGC	3465	GCCCAGAA GGCTAGCTACAACGA TCAGGGAGC	4751
278	AGUUCUGG G CAUUCUGC	3466	GCGAAATG GGCTAGCTACAACGA CCAGAACT	4752
280	UUCUGGGC A UUUUCGCC	3467	GGGCGAAA GGCTAGCTACAACGA GCCCAGAA	4753
285	GGCAUUC G CCCGGCUC	3468	GAGCCGGG GGCTAGCTACAACGA GAAATGCC	4754
290	UUCGGCCG G CUCGAGGU	3469	ACCTCGAG GGCTAGCTACAACGA CGGGCGAA	4755
297	GGCUUCGAG G UGCAGGAU	3470	ATCCTGCA GGCTAGCTACAACGA CTCGAGCC	4756
299	CUCGAGGU G CAGGAUGC	3471	GCATCCTG GGCTAGCTACAACGA ACCTCGAG	4757
304	GGUGCAGG A UGCAGAGC	3472	GCTCTGCA GGCTAGCTACAACGA CCTGCACC	4758
306	UGCAAGAU G CAGAGCAA	3473	TTGCTCTG GGCTAGCTACAACGA ATCCTGCA	4759
311	GAUGCAGA G CAAGGGUC	3474	GCACCTTG GGCTAGCTACAACGA TCTGCATC	4760
316	AGAGCAAG G UGCUGCUG	3475	CAGCAGCA GGCTAGCTACAACGA CTTGCTCT	4761
318	AGCAAGGU G CUCUGGGC	3476	GCCAGCAG GGCTAGCTACAACGA ACCTTGCT	4762
321	AAGGUGCU G CUGGGCGU	3477	ACGGCCAG GGCTAGCTACAACGA AGCACCTT	4763
325	UGCUUCGUG G CCGUUCGCC	3478	GGCGACGG GGCTAGCTACAACGA CAGCAGCA	4764
328	UGCUUGGCC G UCGCCCCU	3479	CAGGGCGA GGCTAGCTACAACGA GGCCAGCA	4765
331	UGGGCGUC G CCCUGGG	3480	CCACAGGG GGCTAGCTACAACGA GACGGCCA	4766
336	GUCCCCU G UGGCUCUG	3481	CAGAGCCA GGCTAGCTACAACGA AGGGCGAC	4767
339	GCCCCUGUG G CUCUGCGU	3482	ACGCAGAG GGCTAGCTACAACGA CACAGGGC	4768
344	GUGGCUCU G CGUGGAGA	3483	TCTCCACG GGCTAGCTACAACGA AGAGCCAC	4769
346	GGCTUCUGC G UGGAGACC	3484	GGTCTCCA GGCTAGCTACAACGA GCAGAGCC	4770
352	GCGUGGGAG A CCCGGGCC	3485	GGCCCCGG GGCTAGCTACAACGA CTCCACGC	4771
358	AGACCCGG G CCCUCUCU	3486	AGAGGGCG GGCTAGCTACAACGA CCGGGTCT	4772
361	CCCGGGCC G CCCUCUGUG	3487	CACAGAGG GGCTAGCTACAACGA GGCCCGGG	4773
367	CGGCCUCU G UGGGUUUG	3488	CAAACCCA GGCTAGCTACAACGA AGAGGCAG	4774
371	CUCUGUGG G UUUGCCUA	3489	TAGGCAAA GGCTAGCTACAACGA CCACAGAG	4775
375	GUGGGUUU G CCUAGUGU	3490	ACACTAGG GGCTAGCTACAACGA AAACCCAC	4776
380	UUUGCCUA G UGUUUCUC	3491	GAGAAACA GGCTAGCTACAACGA TAGGCAAA	4777
382	UGCCUAGU G UUUCUCUU	3492	AAGAGAAA GGCTAGCTACAACGA ACTAGGCA	4778
392	UUCUCUUG A UCUGCCCA	3493	TGGGCAGA GGCTAGCTACAACGA CAAGAGAA	4779
396	CUUCAUCU G CCCAGGCC	3494	AGCCTGGG GGCTAGCTACAACGA AGATCAAG	4780
402	CUGCCCCAG G CUCAGCAU	3495	ATGCTGAG GGCTAGCTACAACGA CTGGGCAG	4781
407	CAGGCCUA G CAUACAAA	3496	TTTGTATG GGCTAGCTACAACGA TGAGCCTG	4782
409	GGCUCAGC A UACAAAAA	3497	TTTTTGTA GGCTAGCTACAACGA GCTGAGCC	4783
411	CUCAGCAU A CAAAAAGA	3498	TCTTTTG GGCTAGCTACAACGA ATGCTGAG	4784
419	ACAAAAAG A CAUACUUA	3499	TAAGTATG GGCTAGCTACAACGA CTTTTTGT	4785
421	AAAAAGAC A UACUUTACA	3500	TGTAAGTA GGCTAGCTACAACGA GTCTTTTT	4786
423	AAAGACAU A CUUACAAU	3501	ATTGTAAG GGCTAGCTACAACGA ATGCTTTT	4787
427	ACAUACUU A CAADUUTAG	3502	CTTAATTG GGCTAGCTACAACGA AAGTATGT	4788
430	UACUUAACA A UUAAGGCCU	3503	AGCCTTAA GGCTAGCTACAACGA TGTAAGTA	4789
436	CAAUUAAG G CUUAAUACA	3504	TGTATTAG GGCTAGCTACAACGA CTTAATTG	4790
440	UAAGGCCUA A UACAACUC	3505	GAGTTGTA GGCTAGCTACAACGA TAGCCTTA	4791
442	AGGCUAUU A CAACUCUU	3506	AAGAGTTG GGCTAGCTACAACGA ATTAGCCT	4792

445	CUAAUACA A CUCUUCAA	3507	TTGAAGAG GGCTAGCTACAAACGA TGTATTAG	4793
454	CUCUUCAA A UUACUUGC	3508	GCAAGTAA GGCTAGCTACAAACGA TTGAAGAG	4794
457	UUCAAAUU A CUUGCCAGG	3509	CCTGCAAG GGCTAGCTACAAACGA AATTGAA	4795
461	AAUACUU G CAGGGGAC	3510	GTCCCCTG GGCTAGCTACAAACGA AAGTAATT	4796
468	UGCAAGGG A CAGAGGGG	3511	TCCCTCTG GGCTAGCTACAAACGA CCCCTGCA	4797
476	ACAGAGGG A CUUGGACU	3512	AGTCCAAG GGCTAGCTACAAACGA CCCTCTGT	4798
482	GGACUUGG A CUGGCCUU	3513	AAAGCCAG GGCTAGCTACAAACGA CCAAGTCC	4799
486	UUGGACUG G CUUUGGCC	3514	GGCCAAAG GGCTAGCTACAAACGA CAGTCCAA	4800
492	UGGCUUUG G CCCAAUAA	3515	TTATTGGG GGCTAGCTACAAACGA CAAAGCCA	4801
497	UUGGCCCA A UAAUCAGA	3516	TCTGATTA GGCTAGCTACAAACGA TGGGCCAA	4802
500	GCCCAAUA A UCAGAGUG	3517	CACTCTGA GGCTAGCTACAAACGA TATTGGC	4803
506	UAAUCAGA G UGGCAGUG	3518	CACTGCCA GGCTAGCTACACAGA TCTGATTA	4804
509	UCAGAGUG G CAGUGAGC	3519	GCTCACTG GGCTAGCTACAAACGA CACTCTGA	4805
512	GAGUGGCA G UGAGCAAA	3520	TTTGCTCA GGCTAGCTACAAACGA TGCCACTC	4806
516	GGCAGUGA G CAAAGGGU	3521	ACCCCTTG GGCTAGCTACAAACGA TCACTGCC	4807
523	ACCAAAAG G UGGAGGUG	3522	CACCTCCA GGCTAGCTACAAACGA CCTTTGCT	4808
529	GGGUGGAG G UGACUGAG	3523	CTCAGTCA GGCTAGCTACAAACGA CTCCACCC	4809
532	UGGAGGUG A CUGAGUGC	3524	GCACTCAG GGCTAGCTACAAACGA CACCTCCA	4810
537	GUGACUGA G UCCAGCGA	3525	TCGCTGCA GGCTAGCTACAAACGA TCAGTCAC	4811
539	GACUGAGU G CAGCGAUG	3526	CATCGCTG GGCTAGCTACAAACGA ACTCAGTC	4812
542	UGAGUGCA G CGAUGGCC	3527	GGCCATCG GGCTAGCTACAAACGA TGCACTCA	4813
545	GUGCAGCG A UGGCCUCU	3528	AGAGGCCA GGCTAGCTACAAACGA CGCTGCAC	4814
548	CAGCGAUG G CCUCUUCU	3529	AGAAGAGG GGCTAGCTACAAACGA CATCGCTG	4815
557	CCUCUUCU G UAAGACAC	3530	GTGTCTTA GGCTAGCTACAAACGA AGAAGAGG	4816
562	UCUGUAAG A CACUCACA	3531	TGTGACTG GGCTAGCTACAAACGA CTTACAGA	4817
564	UGUAAGAC A CUCACAAU	3532	ATTGTGAG GGCTAGCTACAAACGA GTCTTACA	4818
568	AGACACUC A CAUUCCA	3533	TGGAATTG GGCTAGCTACAAACGA GAGTGTCT	4819
571	CACUCACA A UUCAAAAA	3534	TTTTGGAA GGCTAGCTACAAACGA TGTGAGTG	4820
580	UUCAAAAA G UGAUCGGA	3535	TCCGATCA GGCTAGCTACAAACGA TTTTGAA	4821
583	AAAAGUG A UCGGAAAU	3536	ATTCCGA GGCTAGCTACAAACGA CACTTTG	4822
590	GAUCGGAA A UGACACUG	3537	CACTGTCA GGCTAGCTACAAACGA TTCCGATC	4823
593	CGGAAAUG A CACUGGAG	3538	CTCCAGTG GGCTAGCTACAAACGA CATTTCG	4824
595	GAAAUGAC A CUGGAGCC	3539	GGCTCCAG GGCTAGCTACAAACGA GTCATTTC	4825
601	ACACUGGA G CCUACAAAG	3540	CTTGTAGG GGCTAGCTACAAACGA TCCAGTGT	4826
605	UGGAGGCCU A CAAGUGCU	3541	AGCACTTG GGCTAGCTACAAACGA AGGCTCCA	4827
609	GCCUACAA G UGCUUCUA	3542	TAGAAGCA GGCTAGCTACAAACGA TTGTAGGC	4828
611	CUACAAAG U CUUCUACC	3543	GGTAGAAG GGCTAGCTACAAACGA ACTTGTAG	4829
617	GUGCUUCU A CCGGGAAA	3544	TTTCCCCG GGCTAGCTACAAACGA AGAACAC	4830
625	ACCGGGAA A CUGACUUG	3545	CAAGTCAG GGCTAGCTACAAACGA TTCCCCGT	4831
629	GGAAACUG A CUUGGCCU	3546	AGGCCAAG GGCTAGCTACAAACGA CAGTTTCC	4832
634	CUGACUUG G CCUCGGUC	3547	GACCGAGG GGCTAGCTACAAACGA CAAGTCAG	4833
640	UGGCCUCG G UCAUUUAU	3548	ATAAAATGA GGCTAGCTACAAACGA CGAGGCCA	4834
643	CCUCGGUC A UUUAUUGUC	3549	GACATAAA GGCTAGCTACAAACGA GACCGAGG	4835
647	GGUCAUUU A UGUCAUAG	3550	CATAGACA GGCTAGCTACAAACGA AAATGACC	4836
649	UCAUUUAU G UCUAUGUU	3551	AACATAGA GGCTAGCTACAAACGA ATAAATGA	4837
653	UUUAUGUCU A UGUUCAAG	3552	CTTGAACA GGCTAGCTACAAACGA AGACATAA	4838
655	AUGUCUAI G UUCAAGAU	3553	ATCTTGAA GGCTAGCTACAAACGA ATAGACAT	4839
662	UGUUCAAG A UUACAGAU	3554	ATCTGTAA GGCTAGCTACAAACGA CTTGAACA	4840
665	UCAAGAUU A CAGAUUCU	3555	GAGATCTG GGCTAGCTACAAACGA AATCTTGA	4841
669	GAUUACAG A UCUCCAUU	3556	AATGGAGA GGCTAGCTACAAACGA CTGTAATC	4842
675	AGAUUCUCC A UUUAUUGC	3557	GCAATAAA GGCTAGCTACAAACGA GGAGATCT	4843
679	CUCCAUU A UUGCUUCU	3558	AGAAGCAA GGCTAGCTACAAACGA AAATGGAG	4844
682	CAUUAUU G CUUCUGUU	3559	AACAGAAG GGCTAGCTACAAACGA AATAAATG	4845

688	UUGCUUCU G UUAGUGAC	3560	GTCACTAA GGCTAGCTACAACGA AGAAGCAA	4846
692	UUCUGUUA G UGACCAAC	3561	GTGGGTCA GGCTAGCTACAACGA TAACAGAA	4847
695	UGUJAGUG A CCAACAAUG	3562	CATGTTGG GGCTAGCTACAACGA CACTAACAA	4848
699	AGUGACCA A CAUGGAGU	3563	ACTCCATC GGCTAGCTACAACGA TGGTCACT	4849
701	UGACCAAC A UGGAGUCG	3564	CGACTCCA GGCTAGCTACAACGA GTGGGTCA	4850
706	AACAUAGGA G UCGGUUAC	3565	GTACACGA GGCTAGCTACAACGA TCCATGTT	4851
709	AUGGAGUC G UGUACAUU	3566	AATGTACA GGCTAGCTACAACGA GACTCCAT	4852
711	GGAGUCGU G UACAUUAC	3567	GTAATGTA GGCTAGCTACAACGA ACGACTCC	4853
713	AGUCGUGU A CAUUAUCUG	3568	CAGTAATG GGCTAGCTACAACGA ACACGACT	4854
715	UCGUGUAC A UUACUGAG	3569	CTCAGTAA GGCTAGCTACAACGA GTACACGA	4855
718	UGUACAUU A CUGAGAAC	3570	GTTCTCAG GGCTAGCTACAACGA AATGTACA	4856
725	UACUGAGA A CAAAAACA	3571	TGTTTTG GGCTAGCTACAACGA TCTCAGTA	4857
731	GAACAAAA A CAAAACUG	3572	CAGTTTG GGCTAGCTACAACGA TTTTGTTC	4858
736	AAAACAAA A CUGUGGUG	3573	CACCAACAG GGCTAGCTACAACGA TTTGTTTT	4859
739	ACAAAACU G UGGUGAUU	3574	AATCACCA GGCTAGCTACAACGA AGTTTTGT	4860
742	AAACUGUG G UGAUUCCA	3575	TGGAATCA GGCTAGCTACAACGA CACAGTTT	4861
745	CUGUGGUG A UUCCAUU	3576	ACATGGAA GGCTAGCTACAACGA CACCACAG	4862
750	GUGAUUCC A UGUCUCGG	3577	CCGAGACCA GGCTAGCTACAACGA GGAATCAC	4863
752	GAUUCCAU G UCUCGGGU	3578	ACCCGAGA GGCTAGCTACAACGA ATGGAATC	4864
759	UGUCUCGG G UCCAUUUC	3579	GAAATGGA GGCTAGCTACAACGA CCGAGACA	4865
763	UCGGGUCC A UUCAAAAU	3580	ATTTGAAA GGCTAGCTACAACGA GGACCCGA	4866
770	CAJUJUCAA A UCUAACCG	3581	CGTTGAGA GGCTAGCTACAACGA TTGAAATG	4867
776	AAAUCUCA A CGUGUCAC	3582	GTGACACG GGCTAGCTACAACGA TGAGATTT	4868
778	AUCUCAAC G UGUCAUU	3583	AAAGTGACA GGCTAGCTACAACGA TTGAGAT	4869
780	CUAACGU G UCACUUUG	3584	CAAAGTGA GGCTAGCTACAACGA ACGTTGAG	4870
783	AACGUGUC A CUUUGUGC	3585	GCACAAAG GGCTAGCTACAACGA GACACGTT	4871
788	GUCAUUU G UGCAAGAU	3586	ATCTTGCA GGCTAGCTACAACGA AAAGTGAC	4872
790	CACUUUGU G CAAGAUAC	3587	GTATCTTG GGCTAGCTACAACGA ACAAGTG	4873
795	UGUGCAAG A UACCCAGA	3588	TCTGGGTA GGCTAGCTACAACGA CTTGCACA	4874
797	UGCAAGAU A CCCGAAAA	3589	TTTCTGGG GGCTAGCTACAACGA ATCTTGCA	4875
810	AAAAAGAG A UUJGUUCC	3590	GGAAACAAA GGCTAGCTACAACGA CTCTTTTC	4876
814	AGAGAUUU G UUCCUGAU	3591	ATCAGGAA GGCTAGCTACAACGA AAATCTCT	4877
821	UGUJUCCUG A UGGUAACCA	3592	TGTTACCA GGCTAGCTACAACGA CAGGAACA	4878
824	UCCUGAUG G UAACAGAA	3593	TTCTGTTA GGCTAGCTACAACGA CATCAGGA	4879
827	UGAUGGUA A CAGAAUUU	3594	AAATTCTG GGCTAGCTACAACGA TACCATCA	4880
832	GUACACAGA A UUCCUGG	3595	CCAGGAAA GGCTAGCTACAACGA TCTGTTAC	4881
842	UCCUGGG A CAGCAAGA	3596	TCTTGCTG GGCTAGCTACAACGA CCCAGGAA	4882
845	CUGGGACCA G CAAGAAGG	3597	CCTTCITG GGCTAGCTACAACGA TGTCCCAG	4883
854	CAAGAAGG G CUUUAUCUA	3598	TAGTAAAG GGCTAGCTACAACGA CCTTCTTG	4884
859	AGGGCUUU A CUAUUCCC	3599	GGGAATAG GGCTAGCTACAACGA AAAGCCCT	4885
862	GUUUUACU A UUCCCAAGC	3600	GCTGGGAA GGCTAGCTACAACGA AGTAAAGC	4886
869	UAUJUCCCA G CUACAUUGA	3601	TCATGTAG GGCTAGCTACAACGA TGGGATA	4887
872	UCCUGCU A CAUGAUCA	3602	TGATCATG GGCTAGCTACAACGA AGCTGGGA	4888
874	CCAGCUAC A UGAUCAGC	3603	GCTGATCA GGCTAGCTACAACGA GTAGCTGG	4889
877	GCUUACAU A UCAGCUAU	3604	ATAGCTGA GGCTAGCTACAACGA CATGTAGC	4890
881	CAUGAUCA G CUAAUGCUG	3605	CAGCATAG GGCTAGCTACAACGA TGATCATG	4891
884	GAUCAGCU A UGCUUGGCA	3606	TGCCAGCA GGCTAGCTACAACGA AGCTGATC	4892
886	UCAGCUAU G CUGGCAUG	3607	CATGCCAG GGCTAGCTACAACGA ATAGCTGA	4893
890	CUAUGCUG G CADGGUCU	3608	AGACCATG GGCTAGCTACAACGA CAGCATAG	4894
892	AUGCUUGGC A UGGCUUUC	3609	GAAGACCA GGCTAGCTACAACGA GCCAGCAT	4895
895	CUGGCAUG G UCUUCUGU	3610	ACAGAAGA GGCTAGCTACAACGA CATGCCAG	4896
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907	UCUGUGAA G CAAAAAUU	3612	AATTTTG GGCTAGCTACAACGA TTCAACAGA	4898

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929	UGAAAGUU A CCAGUCUA	3617	TAGACTGG GGCTAGCTACAACGA AACTTTCA	4903
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944	UAUUAUGU A CAUAGUUG	3622	CAACTATG GGCTAGCTACAACGA ACATAATA	4908
946	UUUAGUAC A UAGUUGUC	3623	GACAACTA GGCTAGCTACAACGA GTACATAA	4909
949	UGUACAUJ A UUGUCGUU	3624	AACGACAA GGCTAGCTACAACGA TATGTACA	4910
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958	UUGUCGUU G UAGGGUAU	3627	ATACCTTA GGCTAGCTACAACGA AACGACAA	4913
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1855	GUACCCUU G UUUAUCCAA	3825	TTGGATAA GGCTAGCTACAACGA AAGGGTAC	5111
1858	CCCUUGUU A UCCAAGCG	3826	CGCTTGGA GGCTAGCTACAACGA AACAAAGGG	5112
1864	UUUAUCCAA G CGGCAAAU	3827	ATTTGCCG GGCTAGCTACAACGA TTGGATAA	5113
1867	UCCAAGCG G CAAAUGUG	3828	CACATTG GGCTAGCTACAACGA CGCTTGGA	5114
1871	AGCGGCAA A UGUGUCAG	3829	CTGACACA GGCTAGCTACAACGA TTGCCGCT	5115
1873	CGGCAAAU G UGUCAGCU	3830	AGCTGACA GGCTAGCTACAACGA ATTTGCCG	5116
1875	GCAAAUGU G UCAGCUUU	3831	AAAGCTGA GGCTAGCTACAACGA ACATTGC	5117
1879	AUGUGUCA G CUUUGUAC	3832	GTACAAAG GGCTAGCTACAACGA TGACACAT	5118
1884	UCAGCUUU G UACAAAUG	3833	CATTTGTA GGCTAGCTACAACGA AAAGCTGA	5119
1886	AGCUUUGU A CAAAUGUG	3834	CACATTG GGCTAGCTACAACGA ACAAAAGCT	5120
1890	UUGUACAA A UGUGAAGC	3835	GCTTCACA GGCTAGCTACAACGA TTGTACAA	5121
1892	GUACAAAU G UGAAGCGG	3836	CCGCTTCA GGCTAGCTACAACGA ATTTGTAC	5122
1897	AAUGUGAA G CGGUCAAC	3837	GTTGACCG GGCTAGCTACAACGA TTCACATT	5123
1900	GUGAAGCG G UCAACAAA	3838	TTTGTGAA GGCTAGCTACAACGA CGCTTCAC	5124
1904	AGCGGUCA A CAAAGUCG	3839	CGACTTTG GGCTAGCTACAACGA TGACCGCT	5125
1909	UCAACAAA G UCGGGAGA	3840	TCTCCCGA GGCTAGCTACAACGA TTTGTGAA	5126
1927	GAGAGAGG G UGAUCUCC	3841	GGAGATCA GGCTAGCTACAACGA CCTCTCTC	5127
1930	AGAGGGUG A UCUCCUUC	3842	GAAGGAGA GGCTAGCTACAACGA CACCTCTC	5128
1940	CUCCUUCC A CGUGACCA	3843	TGGTCACG GGCTAGCTACAACGA GGAAGGAG	5129
1942	CCUUCCAC G UGACCAAG	3844	CCTGGTCA GGCTAGCTACAACGA GTGGAAGG	5130
1945	UCCACGUG A CCAGGGGU	3845	ACCCCTGG GGCTAGCTACAACGA CACGTGGA	5131
1952	GACCAGGG G UCCUGAAA	3846	TTTCAGGA GGCTAGCTACAACGA CCCTGGTC	5132
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1963	CUGAAAUU A CUUUGCAA	3848	TTGCAAAG GGCTAGCTACAACGA AATTTCAG	5134
1968	AUUAUCUU G CAACCUUG	3849	TCAGTTG GGCTAGCTACAACGA AAAGTAAT	5135
1971	ACUUUUGCA A CCUGACAU	3850	ATGTCAGG GGCTAGCTACAACGA TGCAAAGT	5136
1976	GCAACCUG A CAUGCAGC	3851	GTCATGG GGCTAGCTACAACGA CAGGTTGC	5137
1978	AACCUUGAC A UGCAGCCC	3852	GGGCTGCA GGCTAGCTACAACGA GTCAGGTT	5138
1980	CCUGACAU G CAGCCCAC	3853	GTGGGCTG GGCTAGCTACAACGA ATGTCAGG	5139
1983	GACAUGCA G CCCACUGA	3854	TCAGTGGG GGCTAGCTACAACGA TGCATGTC	5140
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2004	GAGAGCCU G UCUUUGUG	3859	CACAAAGA GGCTAGCTACAACGA ACGCTCTC	5145
2010	GUGUCUUU G UGGUGCAC	3860	GTGCAACCA GGCTAGCTACAACGA AAAGACAC	5146
2013	UCUUUUGUG G UGCACUGC	3861	GCAGTGCA GGCTAGCTACAACGA CACAAAGA	5147
2015	UUUGUGGU G CACUGGAG	3862	CTGCAGTG GGCTAGCTACAACGA ACCACAAA	5148
2017	UGUGGUGG A CUGCGAC	3863	GTCTGCGA GGCTAGCTACAACGA GCACCCACA	5149
2020	GGUGCACU G CAGACAGA	3864	TCTGTCTG GGCTAGCTACAACGA AGTGCACC	5150
2024	CACUGCAG A CAGAUCAA	3865	TAGATCTG GGCTAGCTACAACGA CTGCAGTG	5151
2028	GCAGACAG A UCUACGUU	3866	AACCTAGA GGCTAGCTACAACGA CTGCTCTC	5152
2032	ACAGAUCA U CGUUUGAG	3867	CTCAAACG GGCTAGCTACAACGA AGATCTGT	5153
2034	AGAUUCUAC G UUUGAGAA	3868	TTCTCAAA GGCTAGCTACAACGA GTAGATCT	5154
2042	GUUUGAGA A CCUCACAU	3869	ATGTGAGG GGCTAGCTACAACGA TCTCAAAC	5155
2047	AGAACCUU A CAUGGUAC	3870	GTACCATG GGCTAGCTACAACGA GAGGTTCT	5156
2049	AACCUUC A UGUUACAA	3871	TTGTACCA GGCTAGCTACAACGA GTGAGGTT	5157
2052	CUCACAU G UACAAGCU	3872	AGCTTGTA GGCTAGCTACAACGA CATGTGAG	5158
2054	CACAUUGG A CAAGCUUG	3873	CAAGCTTG GGCTAGCTACAACGA ACCATGTG	5159
2058	UGGUACAA G CUUGGCCC	3874	GGGCCAAG GGCTAGCTACAACGA TTGTACCA	5160
2063	CAAGCUUG G CCCACAGC	3875	GCTGTGGG GGCTAGCTACAACGA CAAGCTTG	5161
2067	CUUGGCCC A CAGCCUCU	3876	AGAGGCTG GGCTAGCTACAACGA GGGCCAAG	5162
2070	GGCCACACA G CCUCUGCC	3877	GGCAGAGG GGCTAGCTACAACGA TGTGGGCC	5163

2076	CAGCCUCU G CCAAIUCCA	3878	TGGATTGG GGCTAGCTACAACGA AGAGGCTG	5164
2080	CUCUGCCA A UCCAUGUG	3879	CACATGGA GGCTAGCTACAACGA TGGCAGAG	5165
2084	GCCAUUCC A UGGGGAG	3880	CTCCCACA GGCTAGCTACAACGA GGATTGGC	5166
2086	CAAUCCAU G UGGGAGAG	3881	CTCTCCC GGCTAGCTACAACGA ATGGATTG	5167
2094	GUUCCAGA G UUGCCCAC	3882	GTGGGCAA GGCTAGCTACAACGA TCTCCCAC	5168
2097	GGAGAGUU G CCCACACC	3883	GGTGTGGG GGCTAGCTACAACGA AACTCTCC	5169
2101	AGUUGGCC A CACCUGUU	3884	AACAGGTG GGCTAGCTACAACGA GGGCAACT	5170
2103	UUGCCCAC A CCUGUUUG	3885	CAAACAGG GGCTAGCTACAACGA GTGGCAA	5171
2107	CCACACCU G UUUGCAAG	3886	CTTGCAAA GGCTAGCTACAACGA AGGTGTGG	5172
2111	ACCUGUUU G CAAGAACU	3887	AGTTCTTG GGCTAGCTACAACGA AAACAGGT	5173
2117	UUGCAAGA A CUUUGGAA	3888	TATCCAAG GGCTAGCTACAACGA TCTTGCAA	5174
2123	GAACUUGG A UACUCUUU	3889	AAAGAGTA GGCTAGCTACAACGA CCAAGTTC	5175
2125	ACUUGGAA U CUCUUUGG	3890	CCAAAGAG GGCTAGCTACAACGA ATCCAAGT	5176
2136	CUUUGGAA A UUGAAUGC	3891	GCATTCAA GGCTAGCTACAACGA TTCCAAAG	5177
2141	GAAAUGA A UGCCACCA	3892	TGGTGGCA GGCTAGCTACAACGA TCAATTTC	5178
2143	AAUUGAAU G CCACCAUG	3893	CATGGTGG GGCTAGCTACAACGA ATTCAATT	5179
2146	UGAAUGCC A CCAUGUUC	3894	GAACATGG GGCTAGCTACAACGA GGCATTCA	5180
2149	AUGCCACC A UGUUCUCU	3895	AGAGAACAA GGCTAGCTACAACGA GGTGGCAT	5181
2151	GCCACCAU G UUCUCUAA	3896	TTAGAGAA GGCTAGCTACAACGA ATGGTGGC	5182
2159	GUUCUCUA A UAGCACAA	3897	TTGTGCTA GGCTAGCTACAACGA TAGAGAAC	5183
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2164	CUAAUAGC A CAAAUAGC	3899	GTCATTG GGCTAGCTACAACGA GCTATTAG	5185
2168	UAGCACAA A UGACAUUU	3900	AAATGTCA GGCTAGCTACAACGA TTGTGCTA	5186
2171	CACAAAUG A CAUUUUGA	3901	TCAAAATG GGCTAGCTACAACGA CATTGTG	5187
2173	CAAUAGAC A UUUUGAUC	3902	GATCAAAA GGCTAGCTACAACGA GTCATTG	5188
2179	ACAUUUUG A UCAUGGAG	3903	CTCCATGA GGCTAGCTACAACGA CAAAATGT	5189
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2187	AUCAUUGG A CUUAAGAA	3905	TTCTTAAG GGCTAGCTACAACGA TCCATGAT	5191
2195	GCUUAAGA A UGCAUCCU	3906	AGGATGCA GGCTAGCTACAACGA TCTTAAGC	5192
2197	UUAAGAAU G CAUCCUUG	3907	CAAGGATG GGCTAGCTACAACGA ATTCTTAA	5193
2199	AAGAAUGC A UCCUUGCA	3908	TGCAAGGA GGCTAGCTACAACGA GCATTCTT	5194
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2219	CCAAGGAG A CUAUGUCU	3911	AGACATAG GGCTAGCTACAACGA CTCCCTTGG	5197
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2224	GAGACUAU G UCUGCCUU	3913	AAGGCAGA GGCTAGCTACAACGA ATAGTCTC	5199
2228	CUAUJUCU G CCUUGCUC	3914	GAGCAAGG GGCTAGCTACAACGA AGACATAG	5200
2233	UCUGCCUU G CUCAAAGAC	3915	GTCTTGAG GGCTAGCTACAACGA AAGGCAGA	5201
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2248	ACAGGAAG A CCAAGAAA	3917	TTTCTTGG GGCTAGCTACAACGA CTTCTGT	5203
2259	AAGAAAAAG A CAUUGGGU	3918	ACGCAATG GGCTAGCTACAACGA CTTTTCTT	5204
2261	GAAAAGAC A UUGCGUGG	3919	CCACGCAA GGCTAGCTACAACGA GTCTTTTC	5205
2264	AAGACAUU G CGUGGUCA	3920	TGACCAACG GGCTAGCTACAACGA AATGTCTT	5206
2266	GACAUUGC G UGGUCAGG	3921	CCTGACCA GGCTAGCTACAACGA GCAATGTC	5207
2269	AUUGCGUG G UCAGGCAG	3922	CTGCCCTGA GGCTAGCTACAACGA CACGCAAT	5208
2274	GUUGUCAG G CAGCUCAC	3923	GTGAGCTG GGCTAGCTACAACGA CTGACCCAC	5209
2277	GUCAAGCA G CUCACAGU	3924	ACTGTGAG GGCTAGCTACAACGA TGCCCTGAC	5210
2281	GGCAGCUC A CAGUCCUA	3925	TAGGACTG GGCTAGCTACAACGA GAGCTGCC	5211
2284	AGCUCACAC G UCCUAGAG	3926	CTCTAGGA GGCTAGCTACAACGA TGTGAGCT	5212
2292	GUCCUAGA G CGUGUGGC	3927	GCCACACG GGCTAGCTACAACGA TCTAGGAC	5213
2294	CCUAGAGC G UGGCAOCC	3928	GTGCCACAA GGCTAGCTACAACGA GCTCTAGG	5214
2296	UAGAGCGU G UGGCAOCC	3929	GGGTGCCA GGCTAGCTACAACGA ACGCTCTA	5215
2299	AGCGUGUG G CACCCRCG	3930	CGTGGGTG GGCTAGCTACAACGA CACACGCT	5216

2301	CGUGUGGC A CCCACGAU	3931	ATCGTGGG GGCTAGCTACAACGA GCCACACG	5217
2305	UGGCACCC A CGAUCACA	3932	TGTGATCG GGCTAGCTACAACGA GGGTGCCA	5218
2308	CACCCACG A UCACAGGA	3933	TCCCTGTGA GGCTAGCTACAACGA CGTGGGTG	5219
2311	CCACGAUC A CAGGAAAC	3934	GTTTCCTG GGCTAGCTACAACGA GATCGTGG	5220
2318	CACAGGAA A CCUGGAGA	3935	TCTCCAGG GGCTAGCTACAACGA TTCTGTG	5221
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2335	AUCAGACG A CAAGUAUU	3938	AATACTTG GGCTAGCTACAACGA CGTCTGAT	5224
2339	GACGACAA G UAUJUGGG	3939	CCCCAATA GGCTAGCTACAACGA TTGTCGTC	5225
2341	CGACAAGU A UUGGGGAA	3940	TTCCCCAA GGCTAGCTACAACGA ACTTGTG	5226
2351	UGGGGAAA G CAUCGAAG	3941	CTTCGATG GGCTAGCTACAACGA TTCCCCCA	5227
2353	GGGAAAGC A UCGAAGUC	3942	GACTTCGA GGCTAGCTACAACGA GCTTTCCC	5228
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2364	GAAGUCUC A UGCACGGC	3944	GCCGTGCA GGCTAGCTACAACGA GAGACTTC	5230
2366	AGUCUCAU G CACGGCAU	3945	ATGCCGTG GGCTAGCTACAACGA ATGAGACT	5231
2368	UCUCUAUC A CGGCAUCU	3946	AGATGCCG GGCTAGCTACAACGA GCATGAGA	5232
2371	CAUGCACG G CAUCUGGG	3947	CCCATGAG GGCTAGCTACAACGA CGTGCATG	5233
2373	UGCACGGC A UCUGGGAA	3948	TTCCCCAGA GGCTAGCTACAACGA GCCGTGCA	5234
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2395	CUCCACAG A UCAUGUGG	3951	CCACATGA GGCTAGCTACAACGA CTGTGGAG	5237
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2400	CAGAUCAU G UGGUUUAA	3953	TTAAACCA GGCTAGCTACAACGA ATGATCTG	5239
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2419	AUAUAUGAG A CCCUUGUA	3957	TACAAGGG GGCTAGCTACAACGA CTCATTAT	5243
2425	AGACCCUU G UAGAAGAC	3958	GTCTTCTA GGCTAGCTACAACGA AAGGGTCT	5244
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2477	CACTAUCC G CAGAGUGA	3969	TCACTCTG GGCTAGCTACAACGA GGATAGTG	5255
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2495	GAAGGGAGG A CGAAGGCC	3971	GGCCPTCG GGCTAGCTACAACGA CCTCCCTTC	5257
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2507	AGGCCUCU A CACCUUGCC	3973	GGCAGGTG GGCTAGCTACAACGA AGAGGGCT	5259
2509	GCCUCUAC A CCUGCCAG	3974	CTGGCAGG GGCTAGCTACAACGA GTAGAGGC	5260
2513	CUACACCU G CCAGGCAU	3975	ATGCCCTG GGCTAGCTACAACGA AGGTGTAG	5261
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2522	CCAGGCAU G CAGUGUUC	3978	GAACACTG GGCTAGCTACAACGA ATGCCCTGG	5264
2525	GGCAUGCA G UGUUCUUG	3979	CAAGAACCA GGCTAGCTACAACGA TGCGATGCC	5265
2527	CAUGCAGU G UUCUUGGC	3980	GCCAAAGAA GGCTAGCTACAACGA ACTGCATG	5266
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2537	UCUUGGGCU G UGCAAAAG	3982	CTTTTGCA GGCTAGCTACAACGA AGCCAAGA	5268
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2545	GUGCAAAA G UGGAGGCA	3984	TGCCTCCA GGCTAGCTACAACGA TTTTGCAC	5270
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2626	CGGUGAUU G CCAUGUUC	4002	GAACATGG GGCTAGCTACAACGA AATCACCG	5288
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2631	AUUGCCAU G UUCUUCUG	4004	CAGAAGAA GGCTAGCTACAACGA ATGGCAAT	5290
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2653	UUCUUGUC A UCAUCCUA	4008	TAGGATGA GGCTAGCTACAACGA GACAAGAA	5294
2656	UUGUCAUC A UCCUACGG	4009	CCGTAGGA GGCTAGCTACAACGA GATGACAA	5295
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2698	AACUGAAG A CAGGCUAC	4017	GTACCTCG GGCTAGCTACAACGA CTPCAGTT	5303
2702	GAAGACAG G CUACUUGU	4018	ACAAGTAG GGCTAGCTACAACGA CTGCTTTC	5304
2705	GACAGGCCU A CUUGUCCA	4019	TGGACAAG GGCTAGCTACAACGA AGCCTGTC	5305
2709	GGCUACUU G UCCAUUCG	4020	ACGATGGA GGCTAGCTACAACGA AAGTAGCC	5306
2713	ACUUGUCC A UCGUCAUG	4021	CATGACGA GGCTAGCTACAACGA GGACAAGT	5307
2716	UGUCUCAUC G UCAUGGAU	4022	ATCCATGA GGCTAGCTACAACGA GATGGACA	5308
2719	CCAUCGUC A UGGAUCCA	4023	TGGATCCA GGCTAGCTACAACGA GACGATGG	5309
2723	CGUCAUUGG A UCCAGAU	4024	CATCTGG GGCTAGCTACAACGA CCATGACG	5310
2729	GGAUCCAG A UGAACUCC	4025	GGAGFTCA GGCTAGCTACAACGA CTGGATCC	5311
2733	CCAGAUGA A CUCCCCAUU	4026	AATGGGAG GGCTAGCTACAACGA TCATCTGG	5312
2739	GAACUCCC A UUGGAGUA	4027	TCATCCAA GGCTAGCTACAACGA GGGAGTTC	5313
2744	CCCAUUGG A UGAACAUU	4028	AATGTTCA GGCTAGCTACAACGA CCAATGGG	5314
2748	UUGGAUGA A CAUUGUGA	4029	TCACAATG GGCTAGCTACAACGA TCATCCAA	5315
2750	GGAUAGAC A UUGUGAAC	4030	GTTCACAA GGCTAGCTACAACGA GTTCATCC	5316
2753	UGAACAUU G UGAACGAC	4031	GTCGTTCA GGCTAGCTACAACGA AATGTTCA	5317
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2760	UGUGAACG A CUGCCUUA	4033	TAAGGCAG GGCTAGCTACAACGA CGTTACAA	5319
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2768	ACUGCCUU A UGAUGCCA	4035	TGGCATCA GGCTAGCTACAACGA AAGGCCAGT	5321
2771	GCCUUAUG A UGGCAGCA	4036	TGCTGGCA GGCTAGCTACAACGA CATAAGGC	5322

2773	CUUAGAU G CCAGCAA	4037	TTTGTGG GGCTAGCTACAAACGA ATCATAAG	5323
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2781	GCCAGCAA A UGGGAAUU	4039	AATTCCCA GGCTAGCTACAAACGA TTGCTGGC	5325
2787	AAAUGGGA A UUCCCCAG	4040	CTGGGAA GGCTAGCTACAAACGA TCCCATT	5326
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2813	GAAGCUAG G UAAGCCUC	4044	GAGGCTTA GGCTAGCTACAAACGA CTAGCTTC	5330
2817	CUAGGUAA G CCUCUUGG	4045	CCAAGAGG GGCTAGCTACAAACGA TTACCTAG	5331
2825	GCCCUUUG G CCGUGGUG	4046	CACCACGG GGCTAGCTACAAACGA CAAGAGGC	5332
2828	UCUUGGCC G UGGUGCCU	4047	AGGCACCA GGCTAGCTACAAACGA GGCCAAGA	5333
2831	UGGCCGUG G UGCCUUUG	4048	CAAAGGCA GGCTAGCTACAAACGA CACGGCCA	5334
2833	GCCGUGGU G CCUUUGGC	4049	GCCAAAGG GGCTAGCTACAAACGA ACCACGGC	5335
2840	UGCCUUUG G CCAAGUGA	4050	TCACTTGG GGCTAGCTACAAACGA CAAAGGCA	5336
2845	UUGGCCAA G UGAUUGAA	4051	TTCAATCA GGCTAGCTACAAACGA TTGGCCAA	5337
2848	GCCAGUG A UUGAAGCA	4052	TGCTTCRA GGCTAGCTACAAACGA CACTTGGC	5338
2854	UGAUUGAA G CAGAUGCC	4053	GGCATCTG GGCTAGCTACAAACGA TTCAATCA	5339
2858	UGAACGAG A UGCCUUUG	4054	CAAAGGCA GGCTAGCTACAAACGA CTGCTTCA	5340
2860	AAGCAGAU G CCUUUGGA	4055	TCCAAAGG GGCTAGCTACAAACGA ATCTGTT	5341
2869	CCUUUGGA A UUGACAAG	4056	CTTGTCAA GGCTAGCTACAAACGA TCCAAAGG	5342
2873	UGGAUUG A CAAGACAG	4057	CTGTCITG GGCTAGCTACAAACGA CAATTCCA	5343
2878	UUGACAAG A CAGCACU	4058	AGTTGCTG GGCTAGCTACAAACGA CTTGTCAA	5344
2881	ACAAGACA G CAACUUGC	4059	GCAAGTTG GGCTAGCTACAAACGA TGTCTTGT	5345
2884	AGACAGCA A CUUGCAGG	4060	CCTGCAAG GGCTAGCTACAAACGA TGCTGTCT	5346
2888	ACCAACUU G CAGGACAG	4061	CTGTCCTG GGCTAGCTACAAACGA AAGTTGCT	5347
2893	CUUGCAGG A CAGUAGCA	4062	TGCTACTG GGCTAGCTACAAACGA CCTGCAAG	5348
2896	GCAGGACA G UAGCAGUC	4063	GAATGCTA GGCTAGCTACAAACGA TGCCCTGC	5349
2899	GGACAGUA G CAGUAAA	4064	TTTGACTG GGCTAGCTACAAACGA TACTGTCC	5350
2902	CAGUTAGCA G UCAAAAUG	4065	CATTTTGA GGCTAGCTACAAACGA TGCTACTG	5351
2908	CAGUAAA A UGUUGAAA	4066	TTTCAACA GGCTAGCTACAAACGA TTGACTG	5352
2910	GUCAAAAU G UUGAAAGA	4067	TCTTTCAA GGCTAGCTACAAACGA ATTITGAC	5353
2923	AAGAAGGA G CAACACAC	4068	GTGTGTG GGCTAGCTACAAACGA TCCTTCTT	5354
2926	AAGGAGCA A CACACAGU	4069	ACTGTGTG GGCTAGCTACAAACGA TGCTCCCT	5355
2928	GGAGCAAC A CACAGUGA	4070	TCACTGTG GGCTAGCTACAAACGA GTTGCTCC	5356
2930	AGCAACAC A CAGUGAGC	4071	GCTCACTG GGCTAGCTACAAACGA GTGTTGCT	5357
2933	AACACACA G UGAGCAUC	4072	GATGCTCA GGCTAGCTACAAACGA TGTGTGTT	5358
2937	CACAGUGA G CAUCGAGC	4073	GCTCGATG GGCTAGCTACAAACGA TCACTGTG	5359
2939	CAGUGAGC A UCGAGCUC	4074	GAGCTCGA GGCTAGCTACAAACGA GCTCACTG	5360
2944	AGCAUCGA G CUCUCAUG	4075	CATGAGAG GGCTAGCTACAAACGA TOGATGCT	5361
2950	GACCUCUC A UGUCUGAA	4076	TTCAGACA GGCTAGCTACAAACGA GAGAGCTC	5362
2952	GUUCUCAU G UCUGAACU	4077	AGTTCAAGA GGCTAGCTACAAACGA ATGAGAGC	5363
2958	AUGUCUGA A CUCAGAU	4078	ATCTTGAG GGCTAGCTACAAACGA TCAGACAT	5364
2965	AACUCAAG A UCCUCAUU	4079	AATGAGGA GGCTAGCTACAAACGA CTTGAGTT	5365
2971	AGAUCCUC A UUCAUAUU	4080	AATATGAA GGCTAGCTACAAACGA GAGGATCT	5366
2975	CCUCAUUC A UAUJUGGUC	4081	GACCAATA GGCTAGCTACAAACGA GAATGAGG	5367
2977	UCAUJUCAU A UGGGUCAC	4082	GTGACCAA GGCTAGCTACAAACGA ATGAATGA	5368
2981	UCAUJUUG G UCACCAUC	4083	GATGGTGA GGCTAGCTACAAACGA CAATATGA	5369
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2987	UGGUCCACC A UCUCAAUG	4085	CATTGAGA GGCTAGCTACAAACGA GGTGACCA	5371
2993	CCAUCUCA A UGUGGUCA	4086	TGACCAACA GGCTAGCTACAAACGA TGAGATGG	5372
2995	AUCUCAAU G UGGUCAAC	4087	GTTGACCA GGCTAGCTACAAACGA ATTGAGAT	5373
2998	UCAAUGUG G UCAACCUC	4088	AAGGTTGA GGCTAGCTACAAACGA CACATTGA	5374
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3017	AGGUGGCC U UACCAAGC	4092	GCTTGGTA GGCTAGCTACAACGA AGGCACCT	5378
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3033	CCAGGAGG G CCACUCAU	4095	ATGAGTGG GGCTAGCTACAACGA CCTCTGG	5381
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3049	UGGUGAUU G UGGAAUUC	4100	GAATTCCA GGCTAGCTACAACGA AATCACCA	5386
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3186	CUGAAACG G CGCUUGGA	4126	TCCAAGCG GGCTAGCTACAACGA CGTTTCAG	5412
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3857	GACUUJUG G CAUJGGAAG	4266	CTTCCATG GGCTAGCTACAACGA TCAAAGTC	5552
3859	CUUUGAGC A UGGAAGAG	4267	CTCTTCCA GGCTAGCTACAACGA GCTCAAAG	5553
3869	GGAAGAGG A UUCUGGAC	4268	GTCCAGAA GGCTAGCTACAACGA CCTCTTCC	5554
3876	GAUUCUGG A CUCUCUCU	4269	AGAGAGAG GGCTAGCTACAACGA CCAGAAC	5555
3885	CUCUCUCU G CCTUACCUC	4270	GAGGTAGG GGCTAGCTACAACGA AGAGAGAG	5556
3889	CUCUGCCU A CCUCACCU	4271	AGGTGAGG GGCTAGCTACAACGA AGGCAGAG	5557
3894	CCUACCUC A CCGUUUUC	4272	GAAACAGG GGCTAGCTACAACGA GAGGTAGG	5558
3898	CCUCACCU G UUUCUCUG	4273	ACAGGAAA GGCTAGCTACAACGA AGGTGAGG	5559
3905	UGUUUCCU G UAUJGAGG	4274	CCTCCATA GGCTAGCTACAACGA AGGAAACA	5560
3907	UUUCCUGU A UGGAGGG	4275	CTCCTCCA GGCTAGCTACAACGA ACAGGAAA	5561
3922	AGGAGGAA G UAUJGAC	4276	GTCACATA GGCTAGCTACAACGA TTCCTCCT	5562
3924	GAGGAAGU A UGUACCCC	4277	GGTCACA GGCTAGCTACAACGA ACTTCCTC	5563
3926	GGAAGUAU G UGACCCCC	4278	TGGGGTCA GGCTAGCTACAACGA ATACTTCC	5564
3929	AGUJAUJUG A CCCAAAU	4279	ATTIGGGG GGCTAGCTACAACGA CACATACT	5565
3936	GACCCCAA A UUCCAUUA	4280	TAATGGAA GGCTAGCTACAACGA TTGGGGTC	5566
3941	CAAAUUC C A UUAIUGACA	4281	TGTCATAA GGCTAGCTACAACGA GGAATTG	5567
3944	AUJUCCAJU A UGACAACA	4282	TGTTGTCA GGCTAGCTACAACGA AATGGAAT	5568
3947	CCAUUJUG A CAACACAG	4283	CTGTGTTG GGCTAGCTACAACGA CATAATGG	5569
3950	UUJUGACA A CACAGCG	4284	CTGCTGTTG GGCTAGCTACAACGA TGTCATAA	5570
3952	AUGACAAAC A CAGCAGG	4285	TCCTGCTG GGCTAGCTACAACGA GTTGTCT	5571
3955	ACAACAC A G CAGGAUC	4286	GATTCCCTG GGCTAGCTACAACGA TGTGTTGT	5572
3961	CAGCAGGA A UCAGUCAG	4287	CTGACTGA GGCTAGCTACAACGA TCCTGCTG	5573
3965	AGGAUCA G UCAGUJUC	4288	GATACTGA GGCTAGCTACAACGA TGATTCT	5574
3969	AUCAGUCA G UAUJUGCA	4289	TGCAGATA GGCTAGCTACAACGA TGACTGAT	5575
3971	CAGUCAGU A UCUGCAGA	4290	TCTGCAGA GGCTAGCTACAACGA ACTGACTG	5576
3975	CAGUJAUU G CGAAACAG	4291	CTGTTCTG GGCTAGCTACAACGA AGATACTG	5577
3980	UCUGCAGA A CAGUJAGC	4292	GCTTACTG GGCTAGCTACAACGA TCTGCAGA	5578
3983	GCAGAAAC A UUAGCGAA	4293	TTCGCTTA GGCTAGCTACAACGA TGTTCTGC	5579
3987	AACAGUAA G CGAAGAG	4294	CTCTTTCG GGCTAGCTACAACGA TTACTGTT	5580
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3999	AAGAGCCG G CCUGUGAG	4296	CTCACAGG GGCTAGCTACAACGA CGGCTCTT	5582
4003	GCCGGCCU G UGAGUGUA	4297	TACACTCA GGCTAGCTACAACGA AGGCCGGC	5583
4007	GCCUGUGA G UGUJAAAAA	4298	TTTTTACA GGCTAGCTACAACGA TCACAGGC	5584
4009	CUGUGAGU G UAAAAACA	4299	TGTTTTTA GGCTAGCTACAACGA ACTCACAG	5585
4015	GUGUAAA A CAUJUGAA	4300	TTCAAATG GGCTAGCTACAACGA TTTTACAC	5586
4017	GUAAAAAC A UUUGAAGA	4301	TCTCAAA GGCTAGCTACAACGA GTTTTAC	5587

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4027	UUGAAGAU A UCCCGUUA	4303	TAACGGGA GGCTAGCTACAACGA ATCTTCAA	5589
4032	GAUAUCCC G UUAGAAGA	4304	TCTCTAA GGCTAGCTACAACGA GGGATATC	5590
4041	UUAGAAGA A CCAGAAGU	4305	ACTTCTGG GGCTAGCTACAACGA TCTTCTAA	5591
4048	AACCAGAA G UAAAAGUA	4306	TACTTTA GGCTAGCTACAACGA TTCTGGTT	5592
4054	AAGUAAAA G UAAUCCCA	4307	TGGGATTA GGCTAGCTACAACGA TTTTACTT	5593
4057	UAAAAGUA A UCCCAAGAU	4308	ATCTGGGA GGCTAGCTACAACGA TACTTTTA	5594
4064	AAUCCCA G UGACAACC	4309	GGTTGTCA GGCTAGCTACAACGA CTGGGATT	5595
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4070	AGAUGACA A CCAGACGG	4311	CCGTCTGG GGCTAGCTACAACGA TGTCATCT	5597
4075	ACAACCAG A CGGACAGU	4312	ACTGTCCG GGCTAGCTACAACGA CTGGTTGT	5598
4079	CCAGACGG A CAGUGGUA	4313	TACCACTG GGCTAGCTACAACGA CCGTCTGG	5599
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4087	ACAGUGGU A UGGUUCUU	4316	AAGAACCA GGCTAGCTACAACGA ACCACTGT	5602
4090	UGGUUAUG G UUCUJUGCC	4317	GGCAAGAA GGCTAGCTACAACGA CATAACCAC	5603
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4151	AUCUUUJUG G UGGAAUGG	4326	CCATTCCA GGCTAGCTACAACGA CAAAAGAT	5612
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4161	GGAAUGGU G CCCAGCAA	4329	TTGCTGGG GGCTAGCTACAACGA ACCATTCC	5615
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4172	CAGCAAAA G CAGGGAGU	4331	ACTCCCTG GGCTAGCTACAACGA TTTTGCTG	5617
4179	AGCAGGGG A UCUGUGGC	4332	GCCACAGA GGCTAGCTACAACGA TCCCTGCT	5618
4183	GGGAGUCU G UGGCAUCU	4333	AGATGCCA GGCTAGCTACAACGA AGACTCCC	5619
4186	AGUCUGUG G CAUCUGAA	4334	TTCAGATG GGCTAGCTACAACGA CACAGACT	5620
4188	UCUGUGGC A UCUGAAGG	4335	CCTTCAGA GGCTAGCTACAACGA GCCACAGA	5621
4196	AUCUGAAG G CUAAACCC	4336	GGTTTGAG GGCTAGCTACAACGA TTTCAGAT	5622
4202	AGGCUCAA A CCAQACAA	4337	TTGCTCTGG GGCTAGCTACAACGA TTGAGCCT	5623
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4227	CAGUCCGG A UAUACACUC	4343	GAGTGATA GGCTAGCTACAACGA CCGGACTG	5629
4229	GUCCGGAU A UCACUCCG	4344	CGGAGTGA GGCTAGCTACAACGA ATCCGGAC	5630
4232	CGGAAUAC A CUCCGAUG	4345	CATCGGGAG GGCTAGCTACAACGA GATATCCG	5631
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4241	CUCCGAUG A CACAGACA	4347	TGTCTGTC GGCTAGCTACAACGA CATCGGAG	5633
4243	CCGAUGAC A CAGACACC	4348	GGTGTCTG GGCTAGCTACAACGA GTCATCGG	5634
4247	UGACACAG A CACCACCG	4349	CGGTGGTG GGCTAGCTACAACGA CTGTGTCA	5635
4249	ACACAGAC A CCACCGUG	4350	CACGGTGG GGCTAGCTACAACGA GTCTGTGT	5636
4252	CAGACACC A CCGGUUAC	4351	GTACACGG GGCTAGCTACAACGA GGTGTCTG	5637
4255	ACACCCAC G UGUACUCC	4352	GGAGTACA GGCTAGCTACAACGA GGTGGTGT	5638
4257	ACCACCGU G UACUCCAG	4353	CTGGAGTA GGCTAGCTACAACGA ACGGTGGT	5639
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4278	GAAGCAGA A CUUUAAA	4357	TTTAAAAG GGCTAGCTACAACGA TCTGCTTC	5643
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4403	ACUCCCGG A CAUCACAU	4381	ATGTGATG GGCTAGCTACAACGA CCGGGAGT	5667
4405	UCCCGGAC A UCACAUCA	4382	TCATGTGA GGCTAGCTACAACGA GTCCGGGA	5668
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4417	CAUGAGAG G UCUGCUCA	4385	TGAGCAGA GGCTAGCTACAACGA CTCTCATG	5671
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4919	CUGCAAAU G CAUUGUGU	4486	ACACAATG GGCTAGCTACAACGA ATTGCGAG	5772
4921	GCAAACUG A UUGUGUUU	4487	AAACACAA GGCTAGCTACAACGA GCATTTGC	5773
4924	AAUGCAUU G UGUUUGCU	4488	ACAAACAA GGCTAGCTACAACGA AATGCATT	5774
4926	UGCAUUGU G UUUGCUCU	4489	AGAGCAAA GGCTAGCTACAACGA ACAATGCA	5775
4930	UUGUGUUU G CUCUGGUG	4490	CACCAAGAG GGCTAGCTACAACGA AAACACAA	5776
4936	UUGCUCUG G UGGAGGUG	4491	CACCTCCA GGCTAGCTACAACGA CAGAGCAA	5777
4942	UGGUGGAG G UGGGCADG	4492	CATGCCCA GGCTAGCTACAACGA CTCCACCA	5778
4946	GGAGGGUGG G CAUGGGGU	4493	ACCCCATG GGCTAGCTACAACGA CCACCTCC	5779
4948	AGGUUGGC A UGGGGUCU	4494	AGACCCCA GGCTAGCTACAACGA GCCCACCT	5780
4953	GGCAUUGG G UCUGUUCU	4495	AAACACAGA GGCTAGCTACAACGA CCCATGCC	5781
4957	UGGGGUCA G UUCUGAAA	4496	TTTCAGAA GGCTAGCTACAACGA AGACCCCA	5782
4965	GUUCUGAA A UGUAAAGG	4497	CCTTTACA GGCTAGCTACAACGA TTCAGAAC	5783
4967	UCUGAAAU G UAAAGGGU	4498	ACCCTTTA GGCTAGCTACAACGA ATTTCAGA	5784
4974	UGUAAAGG G UUCAGACG	4499	CGTCTGAA GGCTAGCTACAACGA CTTTACA	5785
4980	GGGUUCAG A CGGGGUUU	4500	AAACCCCG GGCTAGCTACAACGA CTGAACCC	5786
4985	CAGACGGG G UUUCUGGU	4501	ACCAGAAA GGCTAGCTACAACGA CCCGTCTG	5787
4992	GGUUUCUG G UUUUAGAA	4502	TTCTAAAA GGCTAGCTACAACGA CAGAAACC	5788
5002	UUUAGAAG G UUGCGUGU	4503	ACACGCAA GGCTAGCTACAACGA CTTCTAAA	5789
5005	AGAAGGUU G CGUGUUCU	4504	AGAACACG GGCTAGCTACAACGA AACCTTCT	5790
5007	AAGGUUGC G UGUUCUUC	4505	GAAGAACAA GGCTAGCTACAACGA GCAACCTT	5791
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5018	UUCUUCGA G UGGGGCUA	4507	TAGCCCCA GGCTAGCTACAACGA TCGAAGAA	5793
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5029	GGGCUAAA G UAGAGUUC	4509	GAACTCTA GGCTAGCTACAACGA TTAGCCC	5795
5034	AAAGUAGA G UUCGUUGU	4510	ACAAACGAA GGCTAGCTACAACGA TCTACTTT	5796
5038	UAGAGUUC G UUGUGCUG	4511	CAGCACAA GGCTAGCTACAACGA GAACTCTA	5797
5041	AGUUCGUU G UGCUGUUU	4512	AAACAGCA GGCTAGCTACAACGA AACGAACCT	5798
5043	UUCGUUGU G CUGUUUCU	4513	AGAAACAG GGCTAGCTACAACGA ACAACGAA	5799

5046	GUUGUGCU G UUUCUGAC	4514	GTCAGAAA GGCTAGCTACAACGA AGCACAAAC	5800
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5060	GACUCUUA A UGAGAGUU	4516	AACTCTCA GGCTAGCTACAACGA TAGGAGTC	5802
5066	UAAUGAGA G UUCCUUCC	4517	GGAAGGAA GGCTAGCTACAACGA TCTCATTA	5803
5077	CCUUCAG A CCGUUAGC	4518	GCTAACGG GGCTAGCTACAACGA CTGGAAGG	5804
5080	UCCAGACC G UUAGCUGU	4519	ACAGCTAA GGCTAGCTACAACGA GGTCTGGA	5805
5084	GACCGUUA G CUGUCUCC	4520	GGAGACAG GGCTAGCTACAACGA TAACGGTC	5806
5087	CGUUAGCU G UCUCCUUG	4521	CAAGGAGA GGCTAGCTACAACGA AGCTAACG	5807
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5122	AUGAUGCA G CUCUGGCC	4527	AGCCAGAG GGCTAGCTACAACGA TGCAATCAT	5813
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5135	GGCUCCUU G UCUCCCAG	4529	CTGGGAGA GGCTAGCTACAACGA AAGGAGCC	5815
5144	UCUCCCCG G CUGAUCCU	4530	AGGATCAG GGCTAGCTACAACGA CTGGGAGA	5816
5148	CCAGGCUG A UCCUUUAU	4531	ATAAAGGA GGCTAGCTACAACGA CAGCCTGG	5817
5155	GAUCCUUU A UUCAGAAU	4532	ATTCTGAA GGCTAGCTACAACGA AAAGGATC	5818
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5192	AGCUCAAG G CUCCCUGC	4539	GCAGGGAG GGCTAGCTACAACGA CTTGAGCT	5825
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5227	CUGCACAA A CCAGCUUC	4547	GAAGCTGG GGCTAGCTACAACGA TTGTGCAG	5833
5231	ACAAACCA G CUUCUGGU	4548	ACCCAGAAG GGCTAGCTACAACGA TGGTTTGT	5834
5238	AGCUUCUG G UUUCUUCU	4549	AGAAGAAA GGCTAGCTACAACGA CAGAAGCT	5835
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5256	GAAUGAAU A CCCUCAUA	4552	TATGAGGG GGCTAGCTACAACGA ATTCAATT	5838
5262	AUACCCUC A UAUUCUGUC	4553	GACAGATA GGCTAGCTACAACGA GAGGGTAT	5839
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5268	UCAUAIUCU G UCCUGAUG	4555	CATCAGGA GGCTAGCTACAACGA AGATATGA	5841
5274	CUGUCUG A UGUGAUUA	4556	ATATCACA GGCTAGCTACAACGA CAGGACAG	5842
5276	GUCCUGAU G UGAAUAGU	4557	ACATATCA GGCTAGCTACAACGA ATCAAGGAC	5843
5279	CUGAUGUG A UUAGUCUG	4558	CAGACATA GGCTAGCTACAACGA CACATCAG	5844
5281	GAUGUGAU A UGUCUGAG	4559	CTCAGACA GGCTAGCTACAACGA ATCACATC	5845
5283	UGUGAUAU G UCUGAGAC	4560	GTCTCAGA GGCTAGCTACAACGA ATATCACA	5846
5290	UGUCUGAG A CUGAAUGC	4561	GCATTCA GGCTAGCTACAACGA CTCAGACA	5847
5295	GAGACUGA A UGGGGAG	4562	CTCCCGCA GGCTAGCTACAACGA TCAGTCTC	5848
5297	GACUGAAU G CGGGAGGU	4563	ACCTCCCG GGCTAGCTACAACGA ATTCAAGTC	5849
5304	UGGGGGAG G UUCAAUU	4564	ACATTGAA GGCTAGCTACAACGA CTCCCGCA	5850
5309	GAGGUUCA A UGUGAACG	4565	GCTTCACA GGCTAGCTACAACGA TGAACCTC	5851
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5326	UGUGUGUG G UGUCAAAG	4571	CITTCGACA GGCTAGCTACAACGA CACACACA	5857
5328	UGUGUGGU G UCAGGUUU	4572	AACTTTGA GGCTAGCTACAACGA ACCACACA	5858
5334	GUGUCAAA G UUUCAGGA	4573	TCCTGAAA GGCTAGCTACAACGA TTTGACAC	5859
5346	CAGGAAGG A UUUUACCC	4574	GGGTAAAA GGCTAGCTACAACGA CCTTCCTG	5860
5351	AGGAUUUU A CCCUUUUG	4575	CAAAAGGG GGCTAGCTACAACGA AAAATCCT	5861
5359	ACCCUUUU G UUCUUCCC	4576	GGGAAGAA GGCTAGCTACAACGA AAAAGGGT	5862
5371	UUCCCCCU G UCCCCAAC	4577	GTGGGGGA GGCTAGCTACAACGA AGGGGAA	5863
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5393	CUCACCCC G CAACCCAU	4581	ATGGGTTG GGCTAGCTACAACGA GGGGTGAG	5867
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5469	GAAUGAUU A UUAGCCAG	4598	CTGGCTAA GGCTAGCTACAACGA AATCATTC	5884
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5489	UCAAAAUU A UUUUAUAG	4602	CTATAAAA GGCTAGCTACAACGA AATTGTA	5888
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5506	CCCAAAUU A UAAACAUCU	4606	AGATGTTA GGCTAGCTACAACGA AATTGGG	5892
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5511	AUUAUAAAC A UCUAUUGU	4608	ACAATAGA GGCTAGCTACAACGA GTTATAAT	5894
5515	UAACAUCAU A UUGUAUUA	4609	TAATACAA GGCTAGCTACAACGA AGATGTTA	5895
5518	CAUCUAAU G UAUUUAUU	4610	AAATAATA GGCTAGCTACAACGA AATAGATG	5896
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5523	AUUGUAUU A UUUGACU	4612	AGCTTAAA GGCTAGCTACAACGA AATACAAT	5898
5529	UUAUUUAUAG A CUUUUAAC	4613	GTAAAAAG GGCTAGCTACAACGA CTAATAAA	5899
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5570	UUGCCCUU G UUCUGUCC	4622	GGACAGAA GGCTAGCTACAACGA AAGGGCAA	5908
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5597	AAAAGAAA A UGUGUUUU	4624	AAAACACA GGCTAGCTACAACGA TTTCTTTT	5910
5599	AAGAAAAU G UUUUUUUU	4625	AAAAAAACA GGCTAGCTACAACGA ATTTTCTT	5911
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5651	ACUUAFAAG A CAUGCUAU	4639	ATAGCATG GGCTAGCTACAACGA CTTATACT	5925
5653	UAUAAGAC A UGCUAUGG	4640	CCATAGCA GGCTAGCTACAACGA GTCTTATA	5926
5655	UAAGACAU G CUUAGGCA	4641	TGCCATAG GGCTAGCTACAACGA ATGTCTTA	5927
5658	GACAUGCU A UGGCACAU	4642	ATGTGCCA GGCTAGCTACAACGA AGCATGTC	5928
5661	AUGCUAUG G CACAUUAU	4643	TATATGTG GGCTAGCTACAACGA CATAGCAT	5929
5663	GCUAUGGC A CAUAUUUU	4644	AATATATG GGCTAGCTACAACGA GCCATAGC	5930
5665	UAUGGCAC A UAUUUUUA	4645	TAATATATA GGCTAGCTACAACGA GTGCCATA	5931
5667	UGGCACAU A UAUUUUUA	4646	TATAATA GGCTAGCTACAACGA ATGTGCCA	5932
5669	GCACAUAU A UUUAUJAGU	4647	ACTATAAA GGCTAGCTACAACGA ATATGTGC	5933
5673	AUAUAUUU A UAGUCUGU	4648	ACAGACTA GGCTAGCTACAACGA AAATATAT	5934
5676	UAUUUAUA G UCUGUUUA	4649	TAACACAGA GGCTAGCTACAACGA TATAATA	5935
5680	UAUAGUCU G UUUAUJGU	4650	TACATAAA GGCTAGCTACAACGA AGACTATA	5936
5684	GUCUGUUU A UGUAGAAA	4651	TTTCTACA GGCTAGCTACAACGA AAACAGAC	5937
5686	CUGUUUAU G UAGAAACA	4652	TGTTTCTA GGCTAGCTACAACGA ATAACAG	5938
5692	AUGUAGAA A CAA AUGUA	4653	TACATTG GGCTAGCTACAACGA TTCTACAT	5939
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5698	AAACAAAU G UAAUUAU	4655	ATATATTA GGCTAGCTACAACGA ATTTGTTT	5941
5701	CAA AUGUA A UUAUUAU	4656	TTAATATA GGCTAGCTACAACGA TACATTG	5942
5703	AAUGUAAU A UAUUAAAG	4657	CTTTATA GGCTAGCTACAACGA ATTACATT	5943
5705	UGUAAUUAU A UJAAAGCC	4658	GGCTTTAA GGCTAGCTACAACGA ATATTACA	5944
5711	AUAUAAA G CCUUUAU	4659	ATATAAGG GGCTAGCTACAACGA TTAAATAT	5945
5716	AAAGCCUU A UAUUAAA	4660	ATTATATA GGCTAGCTACAACGA AAGGCTT	5946
5718	AGCCUUUAU A UAUAAUGA	4661	TCATTATA GGCTAGCTACAACGA ATAAGGCT	5947
5720	CCUUUAU A UAAUGAAC	4662	GTTCAUTA GGCTAGCTACAACGA ATATAAGG	5948
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5727	UAUUAUGA A CUUGUAC	4664	GTACAAAG GGCTAGCTACAACGA TCATTATA	5950
5732	UGAACUUU G UACUAAUJC	4665	GAATAGTA GGCTAGCTACAACGA AAAGTCA	5951
5734	AACUUUGU A CUAUUCAC	4666	GTGAATAG GGCTAGCTACAACGA ACAAAAGT	5952
5737	UUUGUACU A UUCACAUU	4667	AATGTGAA GGCTAGCTACAACGA AGTACAAA	5953
5741	UACUAAUC A CAUUUGU	4668	ACAAAATG GGCTAGCTACAACGA GAATAGTA	5954
5743	CUAUUCAC A UUUUGUAU	4669	ATACAAAA GGCTAGCTACAACGA GTGAATAG	5955
5748	CACAUUUU G UAUCAUGUA	4670	TACTGATA GGCTAGCTACAACGA AAAATGTG	5956
5750	CAUUUUGU A UCAGUAIU	4671	AATACTGTA GGCTAGCTACAACGA ACAAAATG	5957
5754	UUGUADCA G UAUUAUGU	4672	ACATAATA GGCTAGCTACAACGA TGATACAA	5958

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5756	GUAUCAGU A UUAUGUAG	4673	CTACATAA GGCTAGCTACAACGA ACTGATAC	5959
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5769	GUAGCAUA A CAAAGGUC	4678	GACCTTG GGCTAGCTACAACGA TATGCTAC	5964
5775	UAACAAAG G UCAUAAAUG	4679	CATTATGA GGCTAGCTACAACGA CTTGTTA	5965
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5781	AGGUCAUA A UGCCUUCA	4681	TGAAAGCA GGCTAGCTACAACGA TATGACCT	5967
5783	GUCAUAAA G CUUUCAGC	4682	GCTGAAAG GGCTAGCTACAACGA ATTATGAC	5968
5790	UGCUUUCA G CAAUUGAU	4683	ATCAATTG GGCTAGCTACAACGA TGAAAGCA	5969
5793	UUUCAGCA A UUGAUGUC	4684	GACATCAA GGCTAGCTACAACGA TGCTGAAA	5970
5797	AGCAAUUG A UGUCAUUU	4685	AAATGACA GGCTAGCTACAACGA CAATTGCT	5971
5799	CAAUUGAU G UCAUUUUU	4686	TAAAATGA GGCTAGCTACAACGA ATCAATTG	5972
5802	UUGAUGUC A UUUUAUUA	4687	TAATAAAA GGCTAGCTACAACGA GACATCAA	5973
5807	GUCAUUUU A UAAAAGAA	4688	TTCTTAA GGCTAGCTACAACGA AAAATGAC	5974
5815	AUAAAAGA A CAUUGAAA	4689	TTTCAATG GGCTAGCTACAACGA TCTTTAAT	5975
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Input Sequence = AF035121. Cut Site = R/Y

Arm Length = 8. Core Sequence = GGCTAGCTACAACGA

AF035121 (Homo sapiens KDR/flk-1 protein mRNA, complete cds.; Acc# AF035121; 5830 bp)

## CLAIMS

1. A compound having Formula II: (SEQ ID NO: 5978)

5' - u, a, c, s, a, au ucU GAg geg aaa gec Gaa Aag aca aB-3'

5       wherein each **a** is 2'-O-methyl adenosine nucleotide, each **g** is a 2'-O-methyl guanosine nucleotide, each **c** is a 2'-O-methyl cytidine nucleotide, each **u** is a 2'-O-methyl uridine nucleotide, each **A** is adenosine, each **G** is guanosine, each **s** individually represents a phosphorothioate internucleotide linkage, U is 2'-deoxy-2'-C-allyl uridine, and **B** is an inverted deoxyabasic moiety.

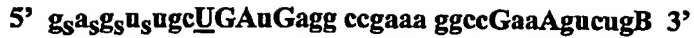
2. A composition comprising the compound of claim 1 and a pharmaceutically acceptable carrier or diluent.
3. A method of administering to a cell the compound of claim 1 comprising contacting said cell with the compound under conditions suitable for said administration.
4. The method of claim 3, wherein said cell is a mammalian cell.
5. The method of claim 3, wherein said cell is a human cell.
6. The method of claim 3, wherein said administration is in the presence of a delivery reagent.
- 20   7. The method of claim 6, wherein said delivery reagent is a lipid.
8. The method of claim 7, wherein said lipid is a cationic lipid.
9. The method of claim 7, wherein said lipid is a phospholipid.
10. The method of claim 6, wherein said delivery reagent is a liposome.
11. A method of administering to a cell the compound of claim 1 in conjunction with one or more other drug comprising contacting said cell

with the compound and the other drug(s) under conditions suitable for said administration.

12. A method of inhibiting ocular angiogenesis in a subject comprising the step of contacting said subject with the compound of claim 1 under conditions suitable for said inhibition.  
5
13. The method of claim 12, wherein said angiogenesis is associated with diabetic retinopathy.
14. The method of claim 12, wherein said angiogenesis is associated with age related diabetic retinopathy.
- 10 15. A method of cleaving RNA comprising a sequence of KDR RNA comprising contacting the compound of claim 1 with said RNA under conditions suitable for the cleavage of said RNA.
16. The method of claim 15, wherein said cleavage is carried out in the presence of a divalent cation.
- 15 17. The method of claim 16, wherein said divalent cation is Mg<sup>2+</sup>.
18. A method of administering to a mammal the compound of claim 1 comprising contacting said mammal with the compound under conditions suitable for said administration.
19. The method of claim 18, wherein said mammal is a human.
- 20 20. The method of claim 18 wherein said administration is in the presence of a delivery reagent.
21. The method of claim 18, wherein said delivery reagent is a lipid.
22. The method of claim 21, wherein said lipid is a cationic lipid.
23. The method of claim 21, wherein said lipid is a phospholipid.
- 25 24. The method of claim 20, wherein said delivery reagent is a liposome.

25. A method for treating a subject having endometriosis, comprising contacting said subject with a nucleic acid molecule that modulates the expression of VEGF, VEGFR1, and/or VEGFR2, under conditions suitable for said treatment.
- 5    26. The method of claim 25, wherein said nucleic acid molecule is an enzymatic nucleic acid molecule.
27. The method of claim 25, wherein said nucleic acid molecule is an antisense nucleic acid molecule.
- 10    28. The method of claim 25, wherein said nucleic acid molecule is a dsRNA nucleic acid molecule.
29. The method of claim 25, wherein said nucleic acid molecule is a nucleic acid aptamer.
30. The method of claim 25, wherein said nucleic acid molecule comprises a sequence having SEQ ID NO: 5977.
- 15    31. The method of claim 26, wherein said enzymatic nucleic acid molecule has an endonuclease activity to cleave RNA encoded by an VEGFR1 and/or VEGFR2 gene.
32. The method of claim 26, wherein said enzymatic nucleic acid molecule is in a hammerhead configuration.
- 20    33. The method of claim 26, wherein said enzymatic nucleic acid molecule is in an Inozyme configuration.
34. The method of claim 26, wherein said enzymatic nucleic acid molecule is in a Zinzyme configuration.
- 25    35. The method of claim 26, wherein said enzymatic nucleic acid molecule is in a DNAzyme configuration.
36. The method of claim 26, wherein said enzymatic nucleic acid molecule is in a G-cleaver configuration.
37. The method of claim 26, wherein said enzymatic nucleic acid molecule is in an Amberzyme configuration.

38. The method of claim 26, wherein said enzymatic nucleic acid molecule is an allozyme.
39. The method of claim 25, wherein said nucleic acid molecule is chemically synthesized.
- 5 40. The method of claim 25, wherein said nucleic acid molecule comprises at least one 2'-sugar modification.
41. The method of claim 25, wherein said nucleic acid molecule comprises at least one nucleic acid base modification.
42. The method of claim 25, wherein said nucleic acid molecule comprises at 10 least one phosphate backbone modification.
43. The method of claim 25, wherein said subject is a human.
44. A method for treating a subject having endometriosis, comprising administering to the subject a nucleic acid molecule that modulates the expression of VEGF, VEGFR1, and/or VEGFR2, under conditions suitable 15 for said treatment.
45. The method of claim 44 wherein said administration is in the presence of a delivery reagent.
46. The method of claim 45, wherein said delivery reagent is a lipid.
47. The method of claim 46, wherein said lipid is a cationic lipid.
- 20 48. The method of claim 46, wherein said lipid is a phospholipid.
49. The method of claim 45, wherein said delivery reagent is a liposome.
50. The method of claim 44, further comprising administering one or more other drug(s).
51. The method of claim 50, wherein said other drug(s) are chosen from GnRH 25 (gonadotropin releasing hormone) agonists, Lupron Depot (Leuproide Acetate), Synarel (naferalin acetate), Zolodex (goserelin acetate), Suprefact (buserelin acetate), Danazol, and oral contraceptives.
52. A compound having Formula I: (SEQ ID NO: 5977)



wherein each **a** is 2'-O-methyl adenosine nucleotide, each **g** is a 2'-O-methyl guanosine nucleotide, each **c** is a 2'-O-methyl cytidine nucleotide, each **u** is a 2'-O-methyl uridine nucleotide, each **A** is adenosine, each **G** is guanosine, each **s** individually represents a phosphorothioate internucleotide linkage, **U** is 2'-deoxy-2'-C-allyl uridine, and **B** is an inverted deoxyabasic moiety.

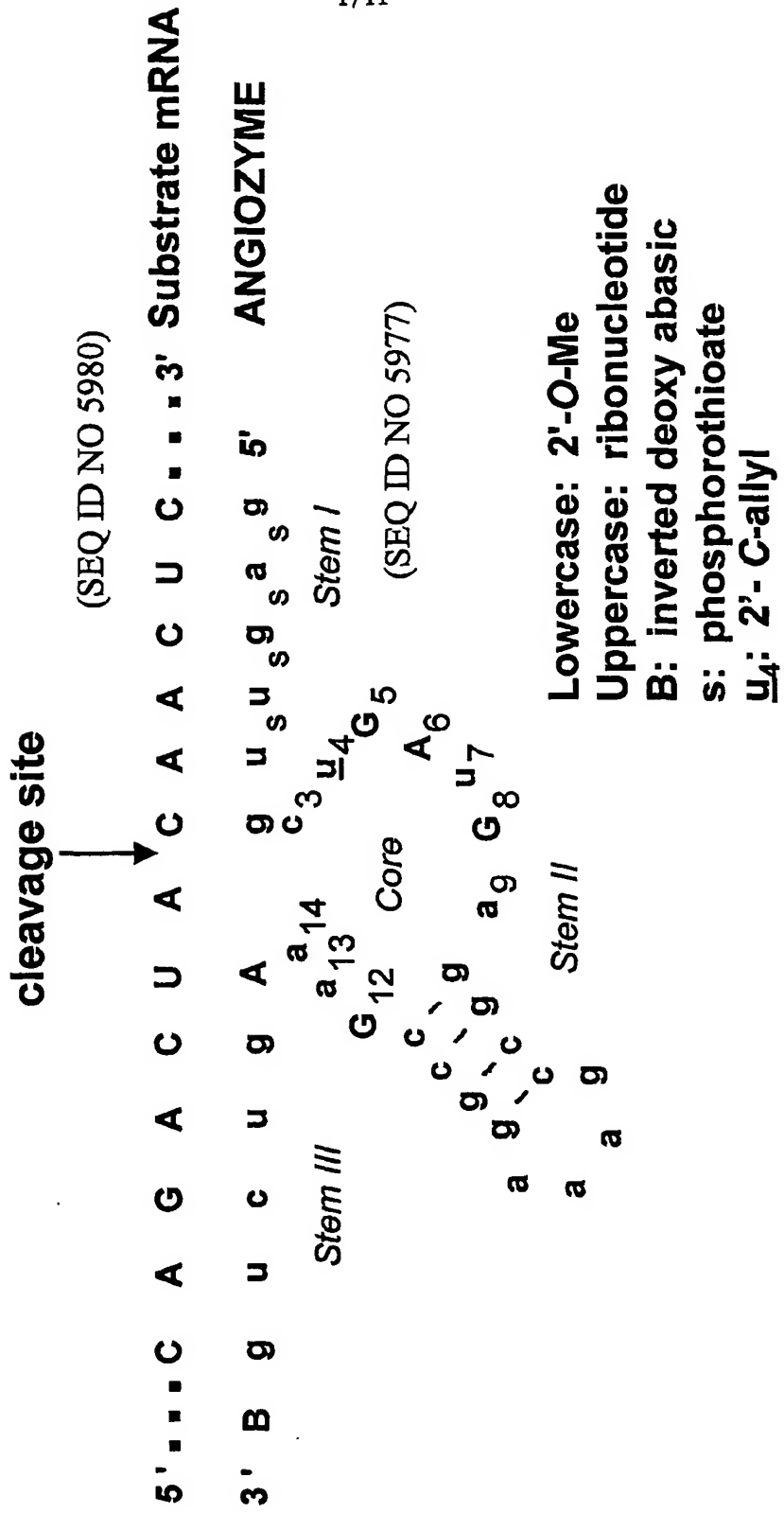
- 5        53. A composition comprising a compound of claim 52 in a pharmaceutically acceptable carrier or diluent.
- 10      54. A method of administering to a cell the compound of claim 52 comprising contacting said cell with the compound under conditions suitable for said administration.
- 15      55. The method of claim 54, wherein said cell is a mammalian cell.
- 16      56. The method of claim 54, wherein said cell is a human cell.
- 20      57. The method of claim 54, wherein said administration is in the presence of a delivery reagent.
- 21      58. The method of claim 57, wherein said delivery reagent is a lipid.
- 22      59. The method of claim 58, wherein said lipid is a cationic lipid.
- 23      60. The method of claim 58, wherein said lipid is a phospholipid.
- 24      61. The method of claim 57, wherein said delivery reagent is a liposome.
- 25      62. A method of administering to a cell the compound of claim 52 in conjunction with a chemotherapeutic agent comprising contacting said cell with the compound and the chemotherapeutic agent under conditions suitable for said administration.
- 26      63. The method of claim 62, wherein said chemotherapeutic agent is 5-fluoro uridine.

64. The method of claim 62, wherein said chemotherapeutic agent is Leucovorin.
65. The method of claim 62, wherein said chemotherapeutic agent is chosen from Irinotecan, CAMPTOSAR®, CPT-11, Camptothecin-11, or Campto.
- 5 66. The method of claim 62, wherein said chemotherapeutic agent is Paclitaxel.
67. The method of claim 62, wherein said chemotherapeutic agent is Carboplatin.
68. A mammalian cell comprising the compound of claim 52..
69. The mammalian cell of claim 68, wherein said mammalian cell is a human cell.
- 10 70. A method of inhibiting angiogenesis in a subject, comprising the step of contacting said subject with the compound of claim 52, under conditions suitable for said inhibition.
71. The method of claim 70, wherein said angiogenesis is tumor angiogenesis.
- 15 72. A method of treatment of a subject having a condition associated with an increased level of VEGF receptor comprising contacting cells of said subject with the compound of claim 52, under conditions suitable for said treatment.
73. The method of claim 72 further comprising the use of one or more drug therapies under conditions suitable for said treatment.
- 20 74. A method of cleaving RNA comprising a sequence of VEGFR1 (flt-1), comprising contacting the compound of claim 52 with said RNA under conditions suitable for the cleavage of said RNA.
75. The method of claim 74, wherein said cleavage is carried out in the presence of a divalent cation.
- 25 76. The method of claim 75, wherein said divalent cation is Mg<sup>2+</sup>.

77. The method of claim 72, wherein said condition is cancer.
78. The method of claim 77, wherein said cancer is breast cancer.
79. The method of claim 77, wherein said cancer is lung cancer.
80. The method of claim 77, wherein said cancer is colorectal cancer.
- 5 81. The method of claim 77, wherein said cancer is renal cancer.
82. The method of claim 77, wherein said cancer is melanoma.
83. The method of claim 77, wherein said cancer is pancreatic cancer.
84. The method of claim 79, wherein said lung cancer is non-small cell lung carcinoma.
- 10 85. The method of claim 81, wherein said renal cancer is renal cell carcinoma.
86. The method of claim 73, wherein said other therapy is 5-fluoro uridine.
87. The method of claim 73, wherein said other therapy is Leucovorin.
88. The method of claim 73, wherein said other therapy is Irinotecan, CAMPTOSAR®, CPT-11, Camptothecin-11, or Campto.
- 15 89. The method of claim 73, wherein said other therapy is Paclitaxel.
90. The method of claim 73, wherein said other therapy is Carboplatin.
91. A method of administering to a mammal the compound of claim 52 comprising contacting said mammal with the compound under conditions suitable for said administration.
- 20 92. The method of claim 91, wherein said mammal is a human.
93. The method of claim 91, wherein said administration is in the presence of a delivery reagent.
94. The method of claim 93, wherein said delivery reagent is a lipid.

95. The method of claim 94, wherein said lipid is a cationic lipid.
96. The method of claim 94, wherein said lipid is a phospholipid.
97. The method of claim 93, wherein said delivery reagent is a liposome.
98. A method of administering to a mammal the compound of claim 52 in conjunction with a chemotherapeutic agent comprising contacting said mammal with the compound and the chemotherapeutic agent under conditions suitable for said administration.  
5
99. The method of claim 98, wherein said chemotherapeutic agent is 5-fluorouridine.
- 10 100. The method of claim 98, wherein said chemotherapeutic agent is Leucovorin.
101. The method of claim 98, wherein said chemotherapeutic agent is Irinotecan, CAMPTOSAR®, CPT-11, Camptothecin-11, or Campto.
102. The method of claim 98, wherein said chemotherapeutic agent is Paclitaxel.
- 15 103. The method of claim 98, wherein said chemotherapeutic agent is Carboplatin.

*Figure 1: Anti-Flt-1 Ribozyme: ANGIOZYME*



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*Inhibition of LLC-HM Primary Tumor Growth Following Systemic ANGIOZYME*

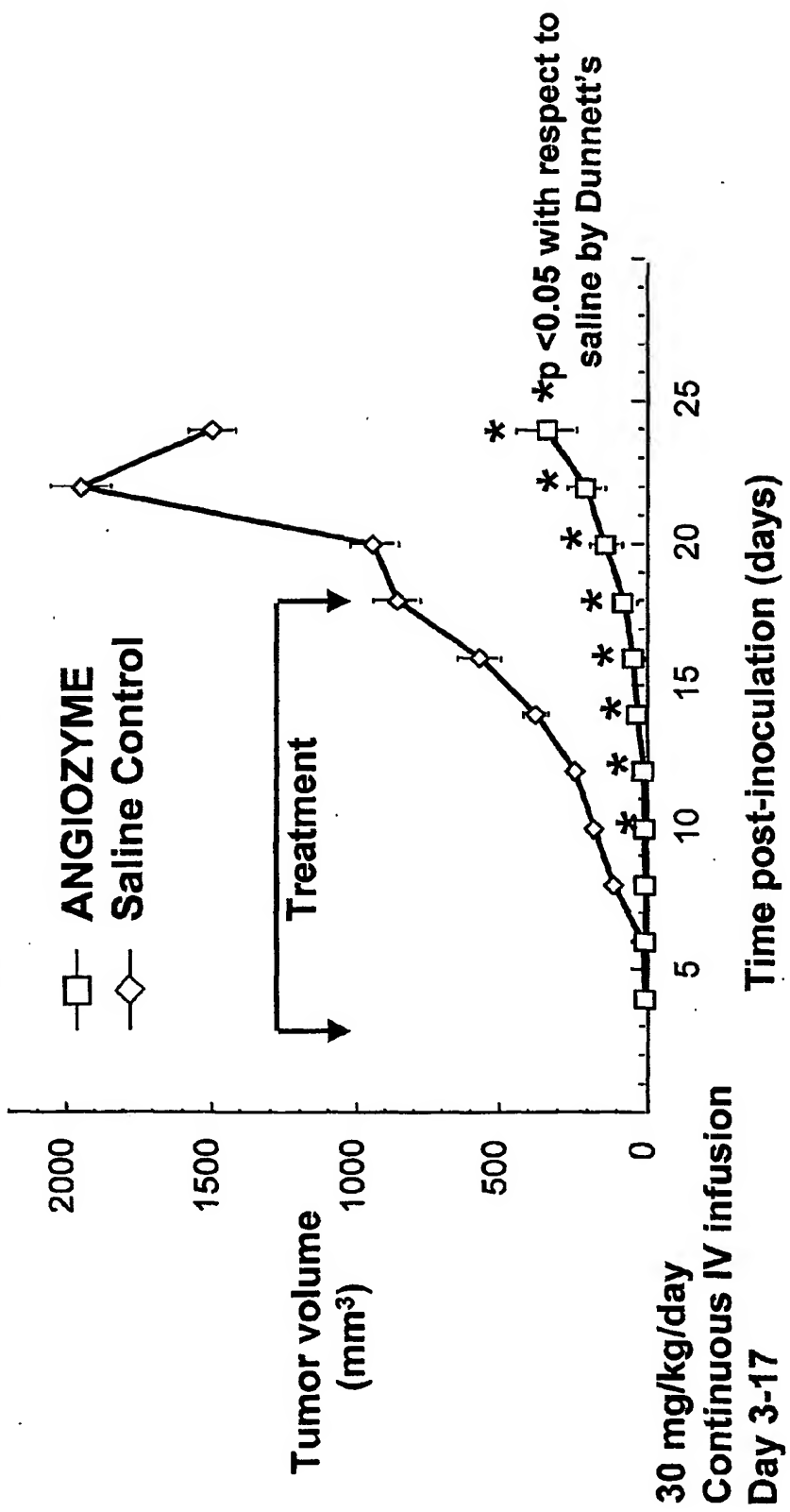
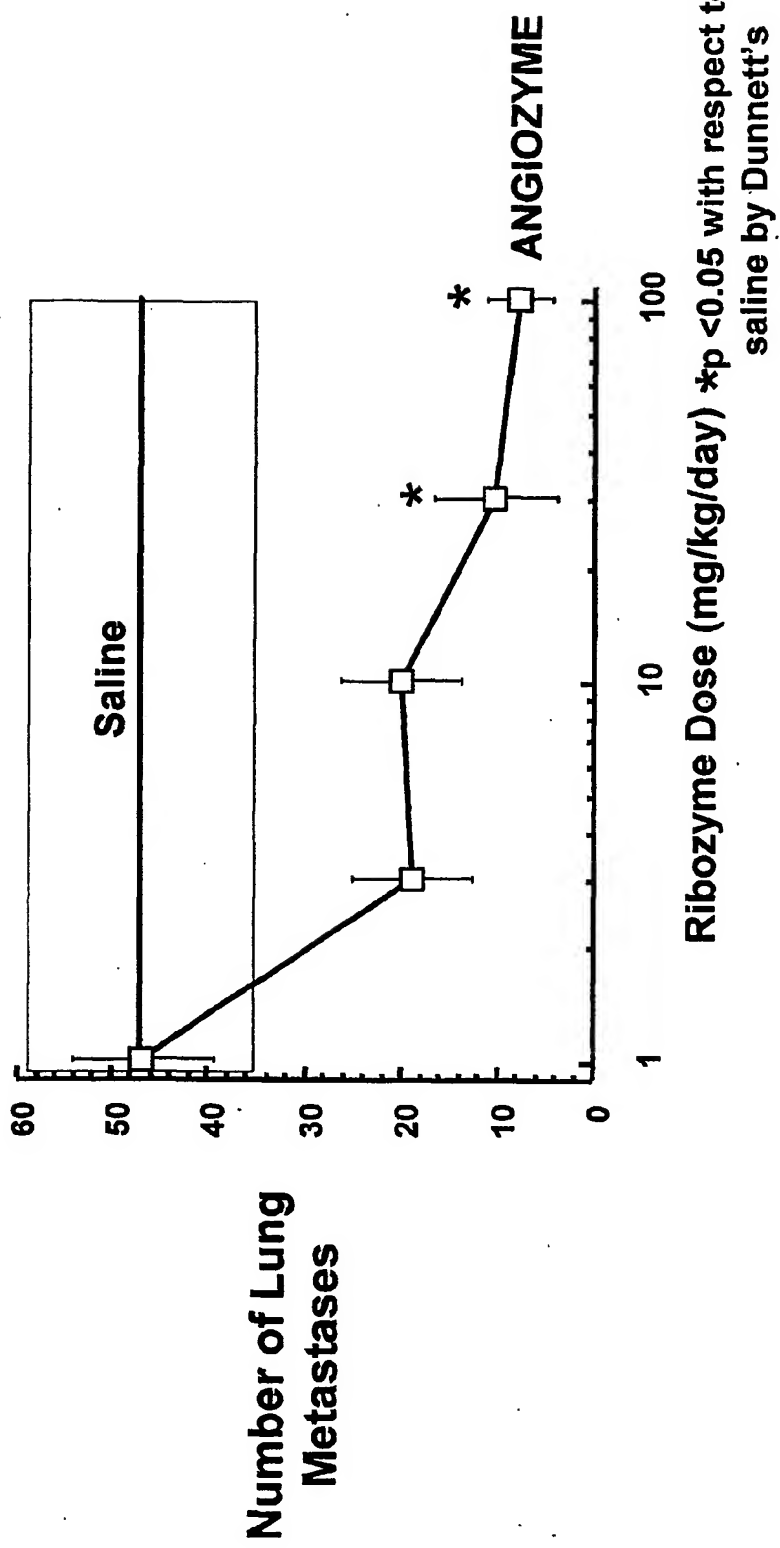


Figure 2

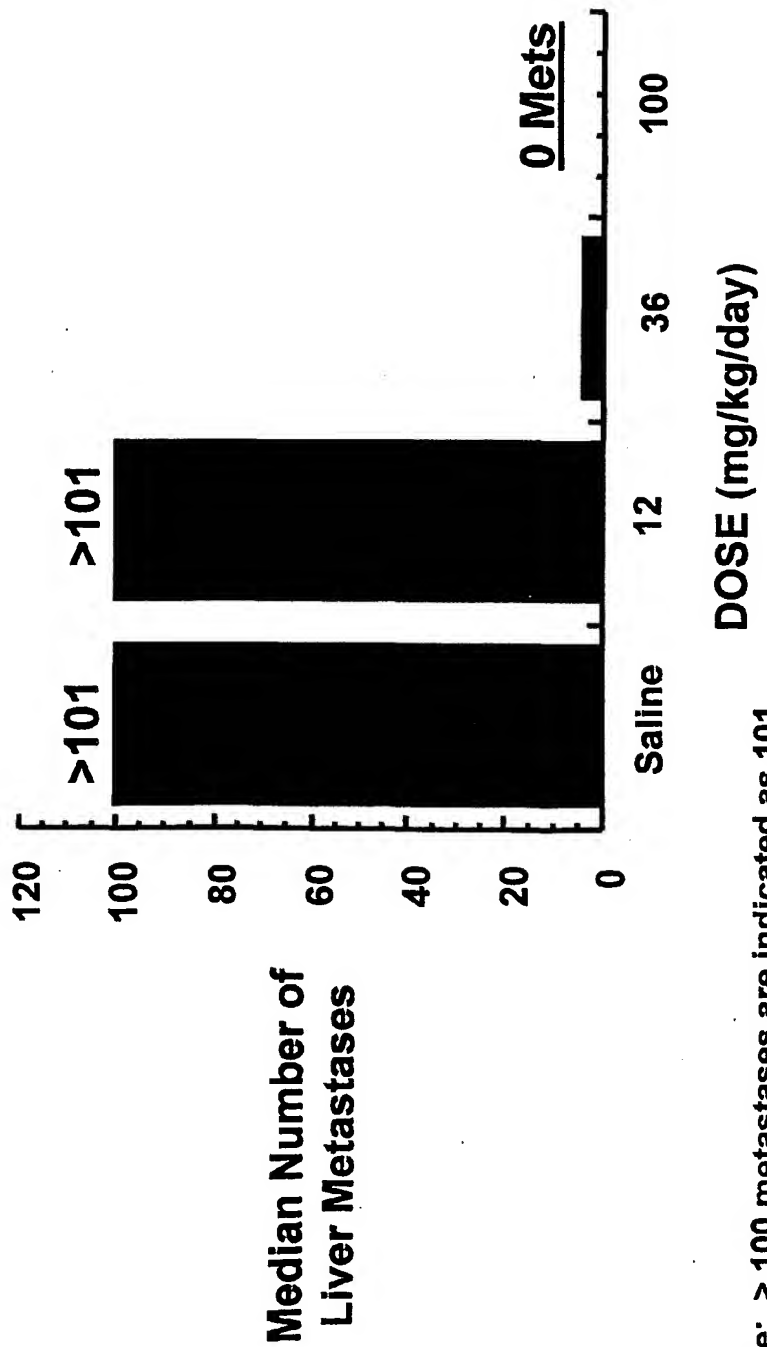
## ANGIOZYME Inhibition of Lung Metastases (LLC-HM Model)



**Figure 3**

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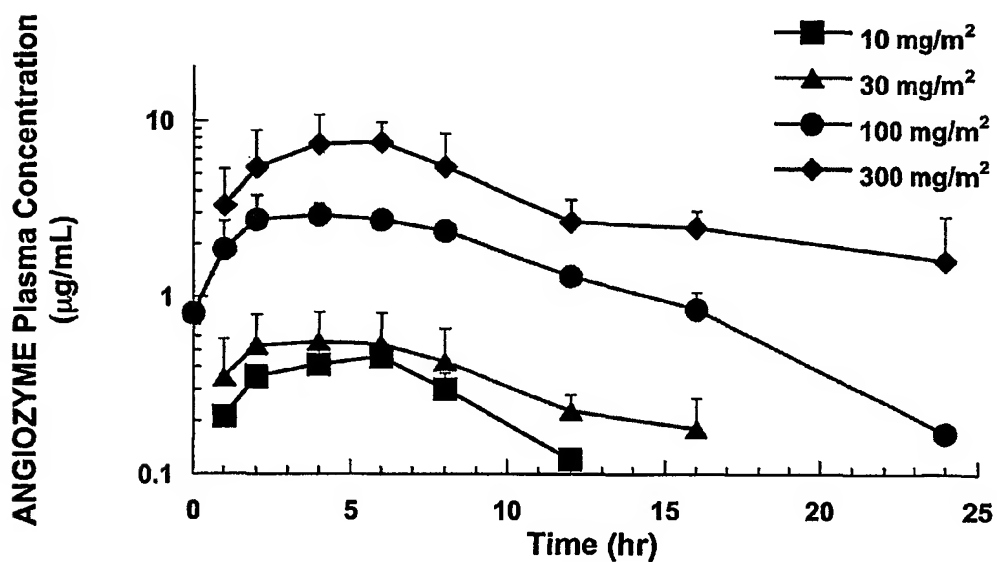
***Effect of ANGIOZYME on Liver Metastases in a Colorectal Cancer Model***



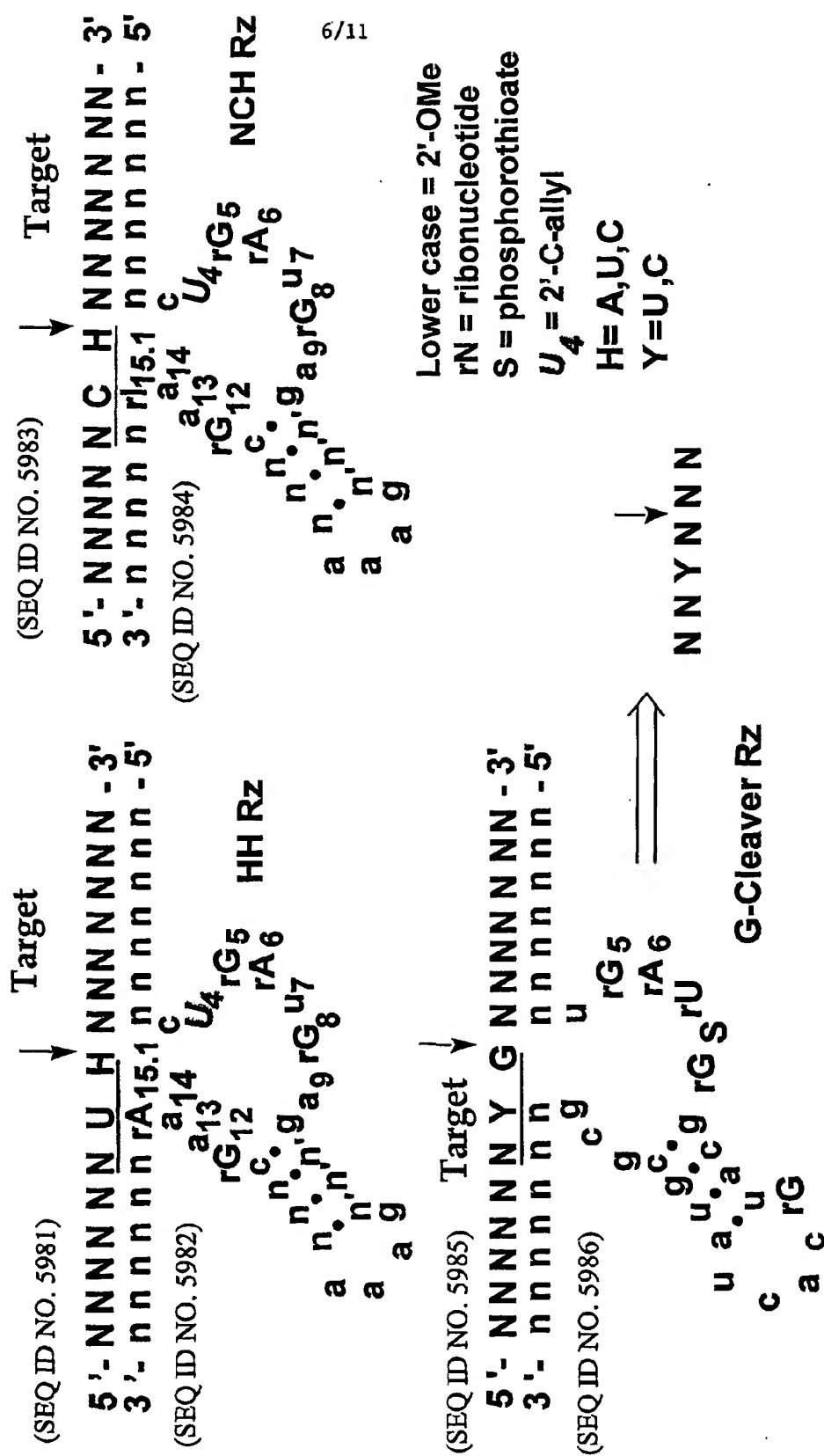
Note: > 100 metastases are indicated as 101.

**Figure 4**

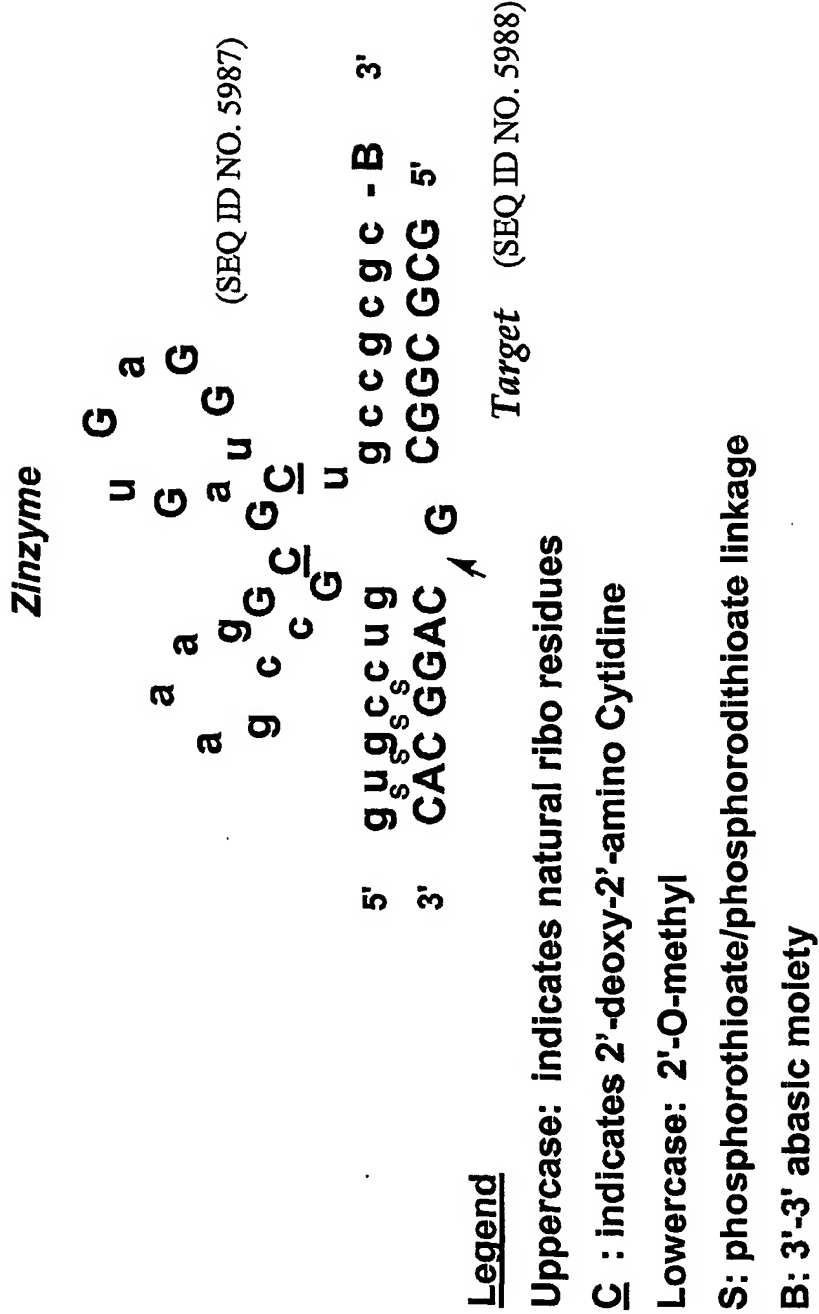
**Figure 5: Plasma concentration profile of ANGIOZYME after a single subcutaneous dose of 10, 30, 100 or 300 mg/m<sup>2</sup>**



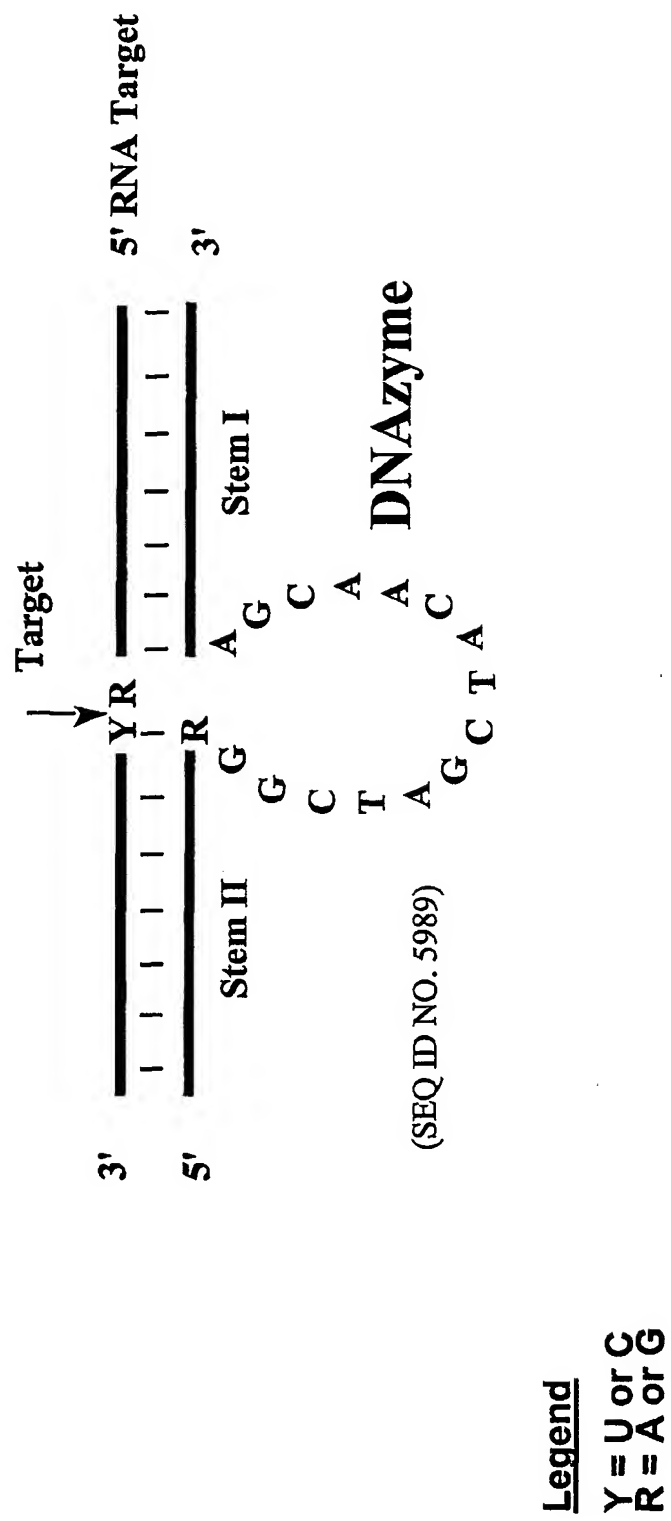
*Figure 6: Examples of Nuclease Stable Ribozyme Motifs*



**Figure 7: Stabilized Zinzyme Ribozyme Motif**

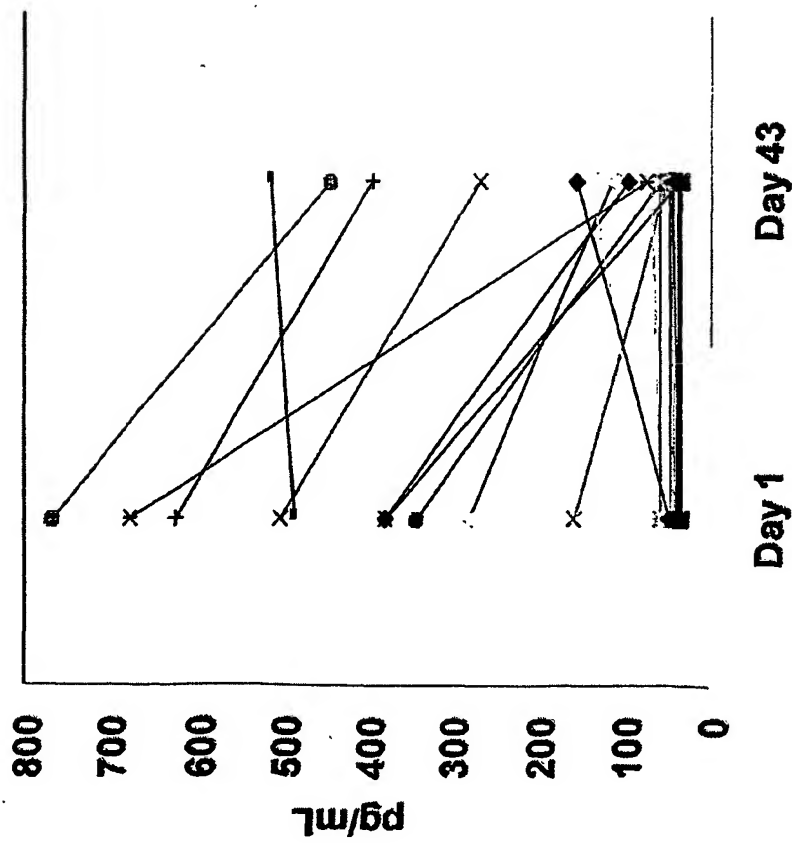


*Figure 8: DNAzyme Motif*

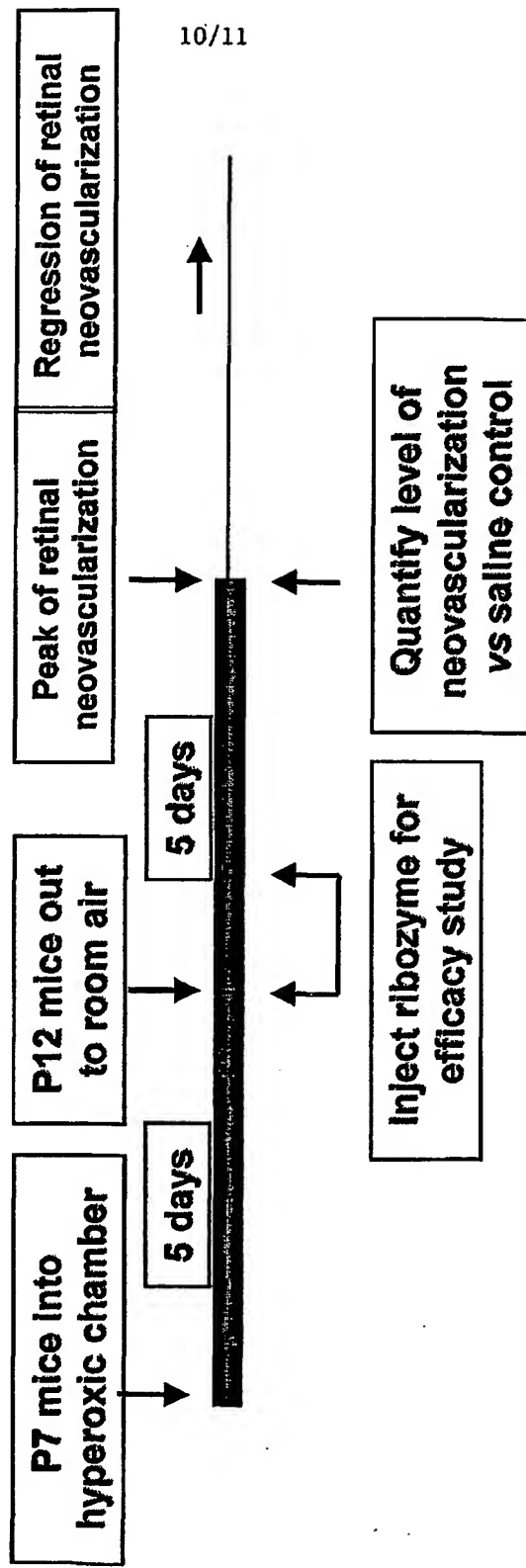


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*Figure 9: Soluble VEGFR1 Reduction*



*Figure 10: Mouse Model of Proliferative Retinopathy*



Note: Peak VEGF levels noted 12 hr after exposure to room air

**Figure 11: RPI.4731 Reduces Hypoxia-Induced Retinal Neovascularization in Neonatal Mice**

